French’s Index of Differential Diagnosis
French’s Index of Differential Diagnosis
An A–Z
16TH EDITION

Mark T. Kinirons BSc Hons MD FRCP FRCP Consultant Physician and Honorary Senior Lecturer, Department of Ageing and Health, Guy’s and St Thomas’ Hospitals, London, UK

Harold Ellis CBE DM MCh FRCS Emeritus Professor of Surgery, Division of Anatomy, Cell and Human Biology, Guy’s, King’s and St Thomas’ School of Biomedical Sciences, London, UK
A Note on Herbert French
(1875–1951)

It might be of interest to readers to learn a little of the original editor of this volume. Herbert French was a scholar at Christ Church, Oxford, and proceeded as a medical student to Guy's Hospital in 1898, with a University Scholarship. He was appointed Assistant Physician at Guy's in 1906 and Full Physician in 1917. He served in the First World War in the Royal Army Medical Corps with the rank of Lieutenant Colonel and was also for many years Physician to the Household of HM George V. French was a prolific writer, and he published An Index of Differential Diagnosis of Main Symptoms in 1912. His ambitious aim was to collect all the symptoms and signs that might arise in the course of disease. He was a man of wide erudition and wrote no less than half of the first edition himself, taking the whole of medicine as his province. The book was an immediate success and was reprinted in the same year and again in 1913 with a second edition appearing in 1917.

H. E.
Contributors

Simon Anderson MD FRCP  
Consultant Gastroenterologist  
Guy’s and St Thomas’ Hospital, London

Michael Baum MD(inc) CPhM FRCS FCRCHon  
Emeritus Professor of Surgery & Visiting  
Professor of Medical Humanities  
University College London

Gerald Carr-White MBBS FRCP PhD  
Consultant Cardiologist and Clinical Lead  
Guy’s and St Thomas’ Hospitals, London

Paul Carroll MD MA MRCPFI FRCP  
Consultant Endocrinologist  
St Thomas’ Hospital, London

Ben Challacombe BSc MS FRCS [Urol]  
Consultant Urological Surgeon  
Guy’s Hospital, London

Duncan Churchill  
Consultant HIV & Tropical Medicine  
Brighton & Sussex University Hospitals, Brighton

Jonathan Compson  
Consultant Orthopaedic and Trauma Surgeon  
The Lister Hospital, London

Reginald Daniel MBBS DO FRCS FRCONTH  
Emeritus Consultant Ophthalmic Surgeon  
Guy’s and St Thomas’ Hospitals, London

Harold Ellis CBE DM MCh FRCS  
Emeritus Professor of Surgery  
Division of Anatomy, Cell and Human Biology, Guy’s,  
King’s and St Thomas’ School of Medicine, London

Toby Garrood FRCP MSc PhD  
Consultant Rheumatologist  
Guy’s and St Thomas’ Hospitals, London

Michael Gleeson MD FRCS FRACS FDS  
Professor of Otolaryngology and Skull Base Surgery  
Guy’s Hospital; and The National Hospital for  
Neurology and Neurosurgery, London

F. A. Graeme Groom  
Consultant Orthopaedic Surgeon  
King’s College Hospital, London

Fred Heatley MA MB BChir(Cantab) FRCS  
Emeritus Professor of Orthopaedics  
King’s College London

Andrew D. Hodgkiss BA MBBS MD FRCPsych DCP  
Consultant Liaison Psychiatrist  
Central & North West London NHS Foundation Trust

Tony Hollingworth MB ChB FRCS(Ed) FRCOG DHMSA PhD MBA  
Consultant in Obstetrics and Gynaecology  
Whipps Cross University Hospital Trust, London

Dipak Kanabar MBBS FRCP FRCPCH  
Consultant Paediatrician  
Guy’s Hospital, London

Mark T. Kinirons BS(Hons) MD FRCPFI FRCP  
Department of Ageing and Health, Guy’s and  
St Thomas’ Hospitals, London

Melvin D. Lobo PhD MRCP  
Consultant Physician  
Bart’s and The London NHS Trust  
Honorary Senior Lecturer  
Bart’s and the London School of Medicine  
and Dentistry, Queen Mary and Westfield College

Jonathan Lucas FRCS FRCS(Ortho&Tr)  
Lead Clinical for Spinal Surgery  
Consultant Spinal Surgeon  
Guys and St Thomas’ NHS Foundation Trust,  
London

Mark McGurk MDS BDS FRCS FDSRCS DLO  
Professor of Oral and Maxillofacial Surgery  
Guy’s and St Thomas’ Hospitals, London

Barry E. Monk MA FRCP  
Consultant Dermatologist  
Dermatology Department, Northampton General  
Hospital, Northampton

James Pattison DM FRCP  
Consultant Nephrologist  
Guy’s Hospital, London

Leandros Vassiliou DDS MD MSc MRCS  
Clinical Fellow in Head and Neck Surgery  
University College London
David J. Werring  FRCP PhD FESO
Professor of Clinical Neurology
Honorary Consultant Neurologist
National Hospital for Neurology and Neurosurgery
and University College Hospital
NIHR CRN: North Thames Clinical Research
Speciality Lead for Stroke

Alex West  MBBS MRCP
Consultant Respiratory Physician
St Thomas’ Hospital, London

IMAGING AND RADIOLOGY THROUGHOUT

Jeremy Rabouhons  BSc (Hons) MBBS (Hons) MRCS FRCR
Specialist Registrar in Radiology, Guy’s and St Thomas’ Hospitals, London
Preface

French's ‘Index’ was first published in 1912. The aim of this volume remains unchanged from the original statement by Herbert French in the first paragraph of his original preface; it is an alphabetic index to help in the differential diagnosis of any condition which may be seen in hospital or general practice. Essentially it is a book for the clinician. With modern transport, regional disease barriers have broken down. Moreover, the time it takes to get anywhere in the world is considerably less than the incubation period of almost all the infectious diseases. So, tropical illnesses are no longer confined to the tropics and one country's epidemic may appear anywhere else in the world in record time. This, together with the massive increase in iatrogenic diseases, makes the art and science of differential diagnosis more interesting than ever before – and vastly more complex too!

The first two editions of this book were edited by Herbert French. Subsequent editors, in turn, were Arthur Douthwaite, his colleague at Guy's Hospital, then Sir Adolphe Abrahams of Westminster Hospital, and then Frank Dudley Hart, also of Westminster. The thirteenth edition had as its editors Professor Ian Bouchier of Edinburgh, the late Peter Fleming of Westminster Hospital, and Harold Ellis. The 14th, 15th and current 16th edition again has Harold Ellis, who is responsible for topics of a ‘surgical’ nature, and he is joined by Mark Kinirons, responsible for the sections on ‘medical’ subjects.

As for the contributors, we have retained a number of old friends and recruited new ones, all chosen carefully for their specialist knowledge and teaching skills. We thank them for their splendid work, although we take full responsibility for the contents of this book. ‘French’ has now been completely revised – many sections are largely rewritten, new ones added, diagnostic methods updated, many old illustrations replaced and others inserted.

The emphasis, however, remains the same – the importance of a careful history, detailed clinical examination and the judicious use of laboratory and imaging investigations in the elucidation of the correct diagnosis.

We hope that this new edition of French's Index will continue to serve the medical profession, both in the United Kingdom and overseas, as it has done now for more than ninety years.

Mark T. Kinirons and Harold Ellis
Professor Heatley is particularly indebted to the late Mr A G Apley, a brilliant teacher who first taught him how to make an orthopaedic diagnosis. Many of the illustrations shown in the orthopaedic sections come from Mr Apley’s slide collection.

The editors would also like to thank the following people for contributing images:

Elizabeth Graham FRCP DO FRCOPHTH
Consultant Medical Ophthalmologist, St Thomas’ Hospital, London

Peter JA Moult MD FRCP
Consultant Physician and Endocrinologist (retired), Whittington Hospital, London

Sheila C Rankin FRCR
Consultant Radiologist, Guys and St Thomas’ Foundation Trust, London
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>ABM</td>
<td>antibasement membrane</td>
</tr>
<tr>
<td>ACL</td>
<td>anterior cruciate ligament</td>
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<tr>
<td>ACS</td>
<td>acute coronary care syndrome</td>
</tr>
<tr>
<td>ACTH</td>
<td>adrenocorticotrophic hormone</td>
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<tr>
<td>ADH</td>
<td>antidiuretic hormone</td>
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<tr>
<td>ADHD</td>
<td>attention-deficit hyperactivity disorder</td>
</tr>
<tr>
<td>AF</td>
<td>atrial fibrillation</td>
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<tr>
<td>AFB</td>
<td>acid-fast bacillus</td>
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<tr>
<td>AIP</td>
<td>acute interstitial pneumonia</td>
</tr>
<tr>
<td>ALS</td>
<td>acid-labile subunit; also amyotrophic lateral sclerosis (ALS)</td>
</tr>
<tr>
<td>AME</td>
<td>apparent mineralocorticoid excess</td>
</tr>
<tr>
<td>AMH</td>
<td>anti-Müllerian hormone</td>
</tr>
<tr>
<td>ANCA</td>
<td>anti-neutrophil cytoplasmic antibody</td>
</tr>
<tr>
<td>ANDI</td>
<td>abnormalities of normal development and involution</td>
</tr>
<tr>
<td>ANUG</td>
<td>acute necrotizing ulcerative gingivitis</td>
</tr>
<tr>
<td>APTT</td>
<td>activated partial thromboplastin time</td>
</tr>
<tr>
<td>ASD</td>
<td>atrial septal defect</td>
</tr>
<tr>
<td>AV</td>
<td>arteriovenous; also atrial-ventricular</td>
</tr>
<tr>
<td>AVM</td>
<td>arteriovenous malformation</td>
</tr>
<tr>
<td>AVNRT</td>
<td>atrial-ventricular node re-entrant tachycardia</td>
</tr>
<tr>
<td>AVRT</td>
<td>atrial-ventricular re-entrant tachycardia</td>
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<tr>
<td>AZT</td>
<td>Zidovudine</td>
</tr>
<tr>
<td>BMI</td>
<td>body mass index</td>
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<tr>
<td>BPE</td>
<td>benign prostatic enlargement</td>
</tr>
<tr>
<td>BPH</td>
<td>benign prostatic hyperplasia</td>
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<tr>
<td>PPFV</td>
<td>benign positional paroxysmal vertigo</td>
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<tr>
<td>BSP</td>
<td>bromsulphthalein</td>
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<tr>
<td>CADASIL</td>
<td>cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy</td>
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<tr>
<td>CAH</td>
<td>congenital adrenal hyperplasia</td>
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<tr>
<td>CAT</td>
<td>computed axial tomography</td>
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<tr>
<td>CBD</td>
<td>common bile duct</td>
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<tr>
<td>CCP</td>
<td>cyclic citrullinated peptides</td>
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<td>CDLE</td>
<td>chronic discoid lupus erythematosus</td>
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<td>CFS</td>
<td>chronic fatigue syndrome</td>
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<td>CDP</td>
<td>chronic inflammatory demyelinating polyneuropathy</td>
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<td>CIN</td>
<td>cervical intra-epithelial neoplasia</td>
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<tr>
<td>CMV</td>
<td>cytomegalovirus</td>
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<tr>
<td>CNS</td>
<td>central nervous system</td>
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<tr>
<td>COPD</td>
<td>chronic obstructive pulmonary disease</td>
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<tr>
<td>CPPD</td>
<td>calcium pyrophosphate dihydrate</td>
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<tr>
<td>CRST</td>
<td>calcinosi, Raynaud’s phenomenon, sclerodactyly, telangiectases (syndrome)</td>
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<tr>
<td>CSD</td>
<td>cortical spreading depression</td>
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<tr>
<td>CSF</td>
<td>cerebrospinal fluid</td>
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<tr>
<td>CSOM</td>
<td>chronic suppurative otitis media</td>
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<tr>
<td>CT</td>
<td>computed tomography</td>
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<td>CTPA</td>
<td>CT pulmonary angiography</td>
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<tr>
<td>CVA</td>
<td>cerebrovascular accident</td>
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<tr>
<td>CXR</td>
<td>chest X-ray</td>
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<tr>
<td>DAT</td>
<td>direct antigen test</td>
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<tr>
<td>DCIS</td>
<td>duct carcinoma-in-situ</td>
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<tr>
<td>DHEA</td>
<td>dehydroepiandrosterone</td>
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<tr>
<td>DHEAS</td>
<td>dehydroepiandrosterone sulphate</td>
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<tr>
<td>DIC</td>
<td>disseminated intravascular coagulation</td>
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<td>DIDMOAD</td>
<td>diabetes insipidus, diabetes mellitus, optic atrophy, deafness (syndrome)</td>
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<td>DISH</td>
<td>diffuse interstitial spinal hyperostosis</td>
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<td>DRPLA</td>
<td>dentatorubral pallidolysian atrophy</td>
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<td>DVT</td>
<td>deep vein thrombosis</td>
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<td>EAA</td>
<td>extrinsic allergic alveolitis</td>
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<td>EBV</td>
<td>Epstein–Barr virus</td>
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<td>ECG</td>
<td>electrocardiography/electrocardiogram</td>
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<td>ECM</td>
<td>erythema chronicum migrans</td>
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<tr>
<td>ECT</td>
<td>electroconvulsive therapy</td>
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<tr>
<td>ED</td>
<td>erectile dysfunction</td>
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<tr>
<td>EEG</td>
<td>electroencephalography/graphic/graph</td>
</tr>
<tr>
<td>ELISA</td>
<td>enzyme-linked immunosorbent assay</td>
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<tr>
<td>EMG</td>
<td>electromyography</td>
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<tr>
<td>EPC</td>
<td>epilepsy partialis continua</td>
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<td>ERCP</td>
<td>endoscopic retrograde cholangiopancreatography</td>
</tr>
<tr>
<td>ESR</td>
<td>erythrocyte sedimentation rate</td>
</tr>
<tr>
<td>EUA</td>
<td>examination under anaesthetic</td>
</tr>
<tr>
<td>EUS</td>
<td>endoscopic ultrasound</td>
</tr>
<tr>
<td>FEV₁</td>
<td>forced expiratory volume in 1 second</td>
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<tr>
<td>FSH</td>
<td>follicle-stimulating hormone</td>
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<tr>
<td>FTA-ABS</td>
<td>fluorescent treponemal antibody absorption</td>
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<tr>
<td>FTD</td>
<td>frontotemporal dementia</td>
</tr>
<tr>
<td>FVC</td>
<td>forced vital capacity</td>
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<tr>
<td>GAVE</td>
<td>gastric antral vascular ectasia</td>
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<tr>
<td>GCA</td>
<td>giant-cell arteritis</td>
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<tr>
<td>GH</td>
<td>growth hormone</td>
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<tr>
<td>GHD</td>
<td>growth hormone deficiency</td>
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<td>GIST</td>
<td>gastrointestinal stromal tumour</td>
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<tr>
<td>GnRH</td>
<td>gonadotrophin-releasing hormone</td>
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</table>
LIST OF ABBREVIATIONS USED

GOR gastro-oesophageal reflux MTP metatarsophalangeal
GTN glyceryl trinitrate NAFL non-alcoholic fatty liver
HAIR-AN hyperandrogenism, insulin resistance, NASH non-alcoholic steatohepatitis
  acanthosis nigricans (syndrome) NHR nuclear hormone receptor
hCG human chorionic gonadotrophin NIPTS noise-induced permanent threshold
HIFU high-intensity focused ultrasound NITTS noise-induced temporary threshold
HMSN hereditary motor–sensory neuropathy shift
HP hypersensitivity pneumonitis NMR nuclear magnetic resonance
HPCR high-pressure chronic retention NPY neuropeptide Y
HPO hypothyroid–pituitary–ovarian axis NSAID non-steroidal anti-inflammatory drug
HPOA hypertrophic pulmonary NSGCT non-seminomatous germ cell tumour
osteopathy
HPV human papilloma virus NSIP non-specific interstitial pneumonia
HRT hormone replacement therapy NSU non-specific urethritis
HSG hysterosalpingogram NTM non-tuberculous mycobacterium
HSV herpes simplex virus NVCD normal variant constitutional delay
HVS hyperventilation syndrome NVSS normal variant short stature
HyCoSy hysterosalpingo contrast sonography OAE otoacoustic emission
IBS irritable bowel syndrome OCD obsessive–compulsive disorder
ICP intracranial pressure OCP oral contraceptive pill
IDO idiopathic detrusor overactivity OGD oesophago-gastro-duodenoscopy,
IGF insulin-like growth factor gastroscopy
IGF1BP insulin-like growth factor binding protein OSA obstructive sleep apnoea
IL-1 interleukin 1 PACI partial anterior circulation infarction
IL-6 interleukin 6 PBC primary biliary cirrhosis
INR International Normalized Ratio PCOS polycystic ovary syndrome
IPT idiopathic thrombocytopenic purpura PE pulmonary embolism
IUCD intra-uterine contraceptive device PEFR peak expiratory flow rate
IVC inferior vena cava PET posiotomion tomography
IVU intravenous urography/gram PID pelvic inflammatory disease
JVP jugular venous pressure PMD post-micturition dribble
KUB kidney–ureter–bladder PMS premenstrual syndrome
LACI lacunar infarction POCI posterior circulation infarction
LDH lactate dehydrogenase POP-Q Pelvic Organ Prolapse Quantification
LFT liver function test PRES posterior reversible encephalopathy
syndrome
LH luteinizing hormone PRIST paper radioimmunosorbent test
LHb lead biventricular RBC red cell
luteinizing hormone-releasing hormone RCI reversible cerebral vasoconstriction
PACI partial anterior circulation infarction
LSD lysergic acid diethylamide PTA post-traumatic amnesia
LVT lead venous occlusion tract PTC percutaneous transhepatic
MALT mucosa-associated lymphoid tissue cholangiogram
MAOI monoamine oxidase inhibitor MD muscular dystrophy PTH parathyroid hormone
MDM mid-diastolic murmur PUL pregnancy of uncertain location
ME myalgic encephalomyelitis PUO pyrexia of unknown origin
MEN multiple endocrine neoplasia RAPD relative afferent pupillary defect
MERRF myoclonic epilepsy with ragged red RAS recurrent apthous stomatitis
fibres RAST radioallergosorbent test
MID multi-infarct disease RCVS reversible cerebral vasoconstriction
MMSE Mini Mental State Examination syndrome
MRI magnetic resonance imaging REM rapid eye movement
MSA-P Parkinsonian variant of multiple system REMI resting energy expenditure
atrophy RSI repetitive strain injury
<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Full Form</th>
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<tr>
<td>RV</td>
<td>residual volume</td>
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<tr>
<td>SCC</td>
<td>squamous cell carcinoma</td>
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<tr>
<td>SeHCAT</td>
<td>selenium homocholic acid taurine</td>
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<tr>
<td>SHBG</td>
<td>sex hormone-binding globulin</td>
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<tr>
<td>SLE</td>
<td>systemic lupus erythematosus</td>
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<tr>
<td>SSRI</td>
<td>selective serotonin reuptake inhibitor</td>
</tr>
<tr>
<td>STI</td>
<td>sexually transmitted infection</td>
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<tr>
<td>SUNCT</td>
<td>short-lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing</td>
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<td>SVC</td>
<td>superior vena cava</td>
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<td>T3</td>
<td>tri-iodothyronine</td>
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<tr>
<td>T4</td>
<td>thyroxine</td>
</tr>
<tr>
<td>TACI</td>
<td>total anterior circulation infarction</td>
</tr>
<tr>
<td>TAR</td>
<td>thrombocytopenia with absent radii</td>
</tr>
<tr>
<td>TCC</td>
<td>transitional cell carcinoma</td>
</tr>
<tr>
<td>TEN</td>
<td>toxic epidermal necrolysis</td>
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<tr>
<td>TGA</td>
<td>transient global amnesia</td>
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<tr>
<td>TIA</td>
<td>transient ischaemic attack</td>
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<tr>
<td>TLC</td>
<td>total lung capacity</td>
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<tr>
<td>TNC</td>
<td>tumour necrosis factor</td>
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<td>TPI</td>
<td>treponemal immobilization (test)</td>
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<td>TSH</td>
<td>thyroid-stimulating hormone</td>
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<td>TURP</td>
<td>transurethral prostatectomy</td>
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<td>UIP</td>
<td>usual interstitial pneumonia</td>
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<td>UMN</td>
<td>upper motor neurone</td>
</tr>
<tr>
<td>UPPP</td>
<td>uvulopharyngoplasty</td>
</tr>
<tr>
<td>VC</td>
<td>vital capacity</td>
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<tr>
<td>VDRL</td>
<td>Venereal Disease Research Laboratory</td>
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<td>VMH</td>
<td>ventromedial hypothalamic nucleus</td>
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<td>VOR</td>
<td>vestibulo-ocular reflex</td>
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<td>VSD</td>
<td>ventricular septal defects</td>
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<tr>
<td>vWF</td>
<td>von Willebrand factor</td>
</tr>
<tr>
<td>XP</td>
<td>xeroderma pigmentosum</td>
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</table>
ABDOMINAL PAIN (GENERAL)

Harold Ellis

(See also ABDOMINAL PAIN, ACUTE, LOCALIZED, p. 3.)

Most abdominal pain is localized, for example that due to a renal stone or biliary stone, acute appendicitis or peptic ulceration. There are, however, a number of causes of generalized abdominal pain, the most common of which are peritonitis and intestinal obstructions.

A list of causes to be considered includes:
1. General peritonitis
2. Tuberculous peritonitis
3. Intestinal obstruction
4. Lead colic (rare)
5. Gastric crises (rare)
6. Abdominal angina
7. Functional abdominal pain
8. General medical diseases:
   • Malaria
   • Porphyria
   • Diabetic ketosis
   • Blood dyscrasias
   • Henoch’s purpura
   • Sickle-cell anaemia
   • Hypercalcaemia

GENERAL PERITONITIS

Peritonitis must be secondary to a lesion that enables some clue in the history to suggest the initiating disease. Thus, the patient with established peritonitis may give a history of onset that indicates acute appendicitis or salpingitis as the source of origin. Where the onset of peritonitis is sudden, one should suspect an acute perforation of a hollow viscus.

The early features depend on the severity and the extent of the peritonitis. Pain is always severe, and typically the patient lies still on its account – in contrast with the restlessness of a patient with abdominal colic. An extensive peritonitis that involves the abdominal aspect of the diaphragm may be accompanied by shoulder-tip pain. Vomiting often occurs early in the course of the disease. The patient is obviously ill, and the temperature frequently elevated. If initially the peritoneal exudate is not purulent, the temperature may be normal.

As the disease progresses, the abdomen becomes distended, signs of free fluid may be detected, and the pulse becomes more rapid and feeble. Vomiting is now effortless and faeculent, and the patient, although still conscious and mentally alert, demonstrates the Hippocratic facies with sunken eyes, pale, cold and sweating skin, and cyanosis of the extremities.

An X-ray of the abdomen in the erect position may reveal free subdiaphragmatic gas in peritonitis due to hollow viscus perforation (e.g. perforated peptic ulcer), but its absence by no means excludes the diagnosis (see Fig. A.1).

The main differential diagnoses are the colics of intestinal obstruction or of ureteric or biliary stone.

Examination of the abdomen demonstrates tenderness, which may be localized to the affected area or is generalized if the peritoneal cavity is extensively involved. There is marked guarding, which again may be localized or generalized, and rebound tenderness is present. The abdomen is silent on auscultation, although sometimes the transmitted sounds of the heart beat and respiration may be detected. Rectally, there is tenderness of the pelvic peritoneum.

Figure A.1 Abdominal radiograph showing the falciform ligament outlined by free intraperitoneal gas (arrowed).
INTESTINAL COLIC

Intraperitoneal haemorrhage, acute pancreatitis, dissection or leakage of an aortic aneurysm, or a basal pneumonia are also important differential diagnoses.

TUBERCULOUS PERITONITIS

In Great Britain, this is now a rare disease. When it is encountered in the UK, the patient is usually an immigrant from a developing country. Usually, there is a feeling of heaviness rather than acute pain. The onset of symptoms is gradual, with abdominal distension, the presence of fluid within the peritoneal cavity, and often the presence of a puckered, thickened omentum, which forms a tumour lying transversely across the middle of the abdomen.

INTESTINAL OBSTRUCTION

This is a common cause of generalized abdominal pain. In peritonitis, there is no periodic rhythm, whereas waves of pain interspersed with periods of complete relief or only a dull ache are typical of obstruction. In contrast to the patients with peritonitis who wish to remain completely still, the victim of intestinal obstruction is restless and rolls about with the spasms of colic. Usually, there are the accompaniments of progressive abdominal distension, absolute constipation, progressive vomiting (which becomes faeculent), and the presence of noisy bowel sounds on auscultation. An X-ray of the abdomen usually reveals multiple fluid levels on the erect film, together with distended loops of gas-filled bowel, which are obvious on the supine radiograph (see Figs C.16 and C.17).

The presence of a scar (or scars) of previous abdominal surgery, performed no matter how long previously, strongly suggests postoperative adhesions or bands as the cause of the obstruction. Careful examination of the hernial orifices – inguinal, femoral and umbilical – is mandatory to diagnose a strangulated external hernia. Surprisingly, the patient may be completely ignorant of its presence. The author has seen a distinguished anaesthetist who correctly diagnosed his own acute bowel obstruction but had not noticed his strangulated inguinal hernia.

LEAD COLIC

Lead colic may cause extremely severe attacks of general abdominal pain. There may be preceding anorexia, constipation and vague abdominal discomfort. The severe pain is usually situated in the lower abdomen and radiates to both groins; it may also sometimes be associated with wrist-drop (due to peripheral neuritis), and occasionally with lead encephalopathy. There may be a blue ‘lead line’ on the gums if oral sepsis is present, due to the precipitation of lead sulphide. Frequently, there is a normocytic hypochromic anaemia with stippling of the red cells (punctuate basophilia). Inquiry about the patient’s occupation may well be the first clue to the diagnosis. Other signs of lead poisoning are considered on p. 229.

GASTRIC CRISIS

Gastric crises in neurosyphilis, although rare, may cause general abdominal pain. The patient has other evidence of tabes dorsalis, with Argyll Robertson pupils, optic atrophy and ptosis, loss of deep sensation (absence of pain on testicular compression or squeezing the Achilles tendon), and loss of ankle and knee jerks. The pain is severe and lasts for many hours or even days. There may be accompanying vomiting, and there may also be rigidity of the abdominal wall. The visceral crisis may be the sole manifestation of tabes. The mere fact that a patient has tabes dorsalis does not, of course, mean that their abdominal pain must necessarily be a gastric crisis. The author has repaired a perforated duodenal ulcer in a patient with all the classic features of well-documented tabes dorsalis.

ABDOMINAL ANGINA

Abdominal angina occurs in elderly patients as a result of progressive atheromatous narrowing of the superior mesenteric artery. Colicky attacks of central abdominal pain occur after meals, and this is followed by diarrhoea. Complete occlusion with infarction of the intestine is often preceded by attacks of this nature. Occlusion of vessels to the small or large intestine – as is seen in a number of vasculopathies such as systemic lupus erythematosus (SLE) or polyarteritis nodosa – may cause generalized abdominal pain and proceed to gangrene, perforation and general peritonitis.

FUNCTIONAL ABDOMINAL PAIN

One of the most difficult problems is the patient (female more often than male) who presents with severe chronic generalized abdominal pains and in whom all clinical, laboratory and radiological tests are negative. Inquiry will often reveal features of depression or the presence of some precipitating factor producing an anxiety state. In some cases, the abdomen is covered with scars of previous laparotomies at which various organs have been repositioned, non-essential viscera removed, and real or imaginary adhesions divided. Some of these patients prove to be drug addicts, others are frank hysterics, and others seek the security of the hospital environment, but in still others the aetiology remains mysterious.
This forms one type of the so-called ‘Munchausen’s syndrome’, described by the late Dr Richard Asher.

**ABDOMINAL PAINS IN GENERAL DISEASE**

Acute abdominal pain may occur in a number of medical conditions not already considered. These include sudden and severe pain complicating malignant malaria, familial Mediterranean fever and cholera, or the pain may accompany uncontrolled diabetes with ketosis, that rare condition known as porphyria and any of the blood dyscrasias; the best examples are Henoch’s purpura in children and the abdominal colic of acute sickle-cell crisis (see p. 54). Bouts of abdominal pain may occur in the hypercalcaemia of hyperparathyroidism.

**ABDOMINAL PAIN, ACUTE, LOCALIZED**

Harold Ellis

A common and extremely important clinical problem is the patient who presents with acute abdominal pain. This may be referred all over the abdominal wall (see ABDOMINAL PAIN (GENERAL), p. 1), but here we shall consider those patients who present pain localized to a particular part of the abdominal cavity.

The causes are legion, and it is a useful exercise to summarize the organs that may be implicated together with the pathological processes pertaining to them so that the clinician can consider the possibilities in a logical manner:

1. **Gastrointestinal**
   - Perforated gastric or duodenal ulcer
   - Perforated gastric carcinoma
   - Acute gastritis (often alcoholic)
   - Irritant poisons
2. **Intestinal**
   - Small-bowel obstruction (adhesions, etc.)
   - Regional ileitis (Crohn’s disease)
   - Intussusception
   - Sigmoid volvulus
   - Acute colonic diverticulitis
   - Large-bowel obstruction due to neoplasm
   - Strangulated external hernia (inguinal, femoral, umbilical)
   - Acute mesenteric occlusion due to arterial embolism or thrombosis or to venous thrombosis
3. **Appendix**
   - Acute appendicitis
4. **Pancreas**
   - Acute pancreatitis
   - Recurrent pancreatitis
   - Pancreatic trauma
5. **Gallbladder and bile ducts**
   - Calculus in the gallbladder or common bile ducts
   - Acute cholecystitis
   - Acute cholangitis
6. **Liver**
   - Trauma
   - Acute hepatitis
   - Malignant disease (primary or secondary)
   - Congestive cardiac failure
7. **Spleen**
   - Trauma
   - Spontaneous rupture (in malaria or infectious mononucleosis)
   - Infarction
8. **Urinary tract**
   - Renal, ureteric or vesical calculus
   - Renal trauma
   - Pyelonephritis
   - Pyonephrosis
9. **Female genitalia**
   - Salpingitis
   - Pyosalpinx
   - Ecstatic pregnancy
   - Torsion of subserous fibroid
   - Red degeneration of fibroid
   - Twisted ovarian cyst
   - Ruptured ovarian cyst
10. **Aorta**
    - Ruptured aneurysm
    - Dissecting aneurysm

In addition to causes from intra-abdominal, retroperitoneal and pelvic organs, it is important to remember that acute localized pain may be referred to the abdomen from other structures:

11. **Central nervous system**
    - Herpes zoster affecting the lower thoracic segments. Posterior nerve root pain (e.g. from prolapsed intervertebral disc or collapsed vertebra from trauma or secondary deposits)
12. **The heart and pericardium**
    - Myocardial infarction
    - Acute pericarditis
13. **Pleura**
    - Acute diaphragmatic pleurisy

Occasionally, patients are seen who are often well known in the Accident and Emergency Department, presenting with simulated acute abdominal pain due to hysteria or malingering.

Patients with acute abdominal pain present one of the most testing trials to the clinician. In the first place, diagnosis is all-important, since a decision has to be made...
be made whether or not the patient requires urgent laparotomy – for example for a perforated peptic ulcer, acute appendicitis or acute intestinal obstruction. The history and examination are often difficult to elicit, particularly in a very ill patient who is in great pain and hardly wishes either to answer a lot of questions or to submit to prolonged examination. Finally, there are very few laboratory or radiological aids to diagnosis. Acute appendicitis, for example, has no specific tests. A raised white blood count suggests intraperitoneal infection, but something like one-quarter of the cases of acute appendicitis have a white blood cell count below 10 000 per mm$^3$. Plain X-rays of the abdomen may indicate free gas when there is a perforated hollow viscus, but this is not invariably so (Fig. A.1). Intestinal obstruction may be revealed by distended loops of bowel on a plain X-ray of the abdomen, but in some 10 per cent of small-bowel obstructions the X-rays are entirely normal, since the distended loops of bowel are filled with fluid only so that the typical gas-distended loops of bowel are not present (see Figs C.16 and C.17).

Ultrasonography of the abdomen may be used to demonstrate distended loops of bowel, fluid collections, gallbladder pathology, the presence of gallstones, a pathological appendix and intussusception. However, accurate diagnosis is heavily observer-dependent and requires the help of an expert ultrasonographer (Fig. A.2). Computed tomography (CT) is also of great assistance, when available (Fig. A.3).

One of the few laboratory investigations that the surgeon relies upon heavily is a raised serum amylase activity. When this is above 1000 units per 100 ml serum, it is almost pathognomic of acute pancreatitis, although every now and then a fulminating case of pancreatitis is seen in which the amylase is not elevated. Unfortunately, more than 200 different assay methods for amylase estimation have been described. Consequently, different hospitals may well have different reference ranges for serum amylase normality. It is therefore essential to know the normal reference range of serum amylase in your own hospital rather than trying to remember values that apply elsewhere. While a very high serum amylase value is typically found in acute pancreatitis and pancreatic trauma, a moderate increase may occur in non-pancreatic acute abdominal disease (e.g. perforated peptic ulcer, intestinal obstruction or infarction). Amylase is cleared from the circulation by the kidneys; anything which interferes with normal renal clearance may therefore also result in a moderate rise in the serum amylase.

Every effort must therefore be made to establish the diagnosis on a careful history and examination. One of the important aspects in the assessment of the acute abdomen is the establishment of a trend. Increasing pain, tenderness, guarding or rigidity indicates that there is some progressive intra-abdominal condition. This is also suggested by a rising pulse rate on hourly or half-hourly observations, and it is also suggested by progressive elevation of the temperature. In a doubtful case, repeated clinical examination – together with sequential recordings of the temperature and pulse – will enable the clinician to decide whether the intra-abdominal condition is either subsiding or progressing.
GENERAL FEATURES
General inspection of the patient is all-important and must never be omitted. The flushed face and coated tongue of acute appendicitis, the agonized expression of the patient with a perforated ulcer, the writhing colic of a patient with ureteric stone, biliary colic or small-bowel obstruction are all most helpful. The skin is inspected for the pallor suggestive of haemorrhage, and for the jaundice that may be associated with biliary colic with a stone impacted at the lower end of the common bile duct. In such a case, there will also be bile pigment that can be detected in the urine.

ABDOMINAL EXAMINATION
The patient must be placed in a good light, and the entire abdomen exposed from the nipples to the knees. The abdomen is inspected. Failure of movement with respiration may suggest an underlying peritoneal irritation. Abdominal distension is present in intestinal obstruction, and visible peristalsis may be seen from rhythmic contractions of the small bowel in these circumstances. Retraction of the abdomen may occur in acute peritonitis so that the abdomen assumes a scaphoid appearance, for example following perforation of a peptic ulcer.

Guarding – a voluntary contraction of the abdominal wall on palpation – denotes underlying inflammatory disease, and this is accompanied by localized tenderness. R rigidity is indicated by an involuntary tightness of the abdominal wall and may be generalized or localized. Localized rigidity over one particular organ suggests local peritoneal involvement, for example in acute appendicitis or acute cholecystitis.

Percussion of the abdomen is useful. Dullness in the flanks suggests the presence of intraperitoneal fluid (e.g. blood in a patient with a ruptured spleen). A resonant distended abdomen is found in obstruction, and loss of liver dullness suggests free gas within the peritoneal cavity in a patient with a ruptured hollow viscus.

In intestinal obstruction, the bowel sounds are increased and have a particular ‘tinkling’ quality. In some cases, borborygmi may be audible without using the stethoscope. A complete absence of bowel sounds suggests peritonitis.

Examination of the abdomen is not complete until the hernial orifices have been carefully inspected and palpated. It is easy enough to miss a small strangulated inguinal, femoral or umbilical hernia that, surprisingly enough, may have been completely overlooked by the patient.

A rectal examination is then performed. In intestinal obstruction, the rectum has a characteristic ‘ballooned’ empty feel, although the exact mechanism of this is unknown. In pelvic peritonitis, there will be tenderness anteriorly in the pouch of Douglas. A tender mass suggests an inflamed or twisted pelvic organ, and this can be confirmed by bimanual vaginal examination.

THE URINE AND SPECIAL INVESTIGATIONS
The presence of blood, protein, pus or bile pigment in the urine may help to distinguish a renal or biliary colic from other causes of intra-abdominal pain. As well as routine testing of a urine specimen, a drop placed under the microscope and viewed with a 1/8th lens (staining is not required) constitutes a useful test. It is the work of a few minutes to see if pus cells or red cells are obvious. In obscure cases of abdominal pain, the urine should be examined for porphyrins to exclude porphyria, particularly when the attack appears to have been precipitated by barbiturates.

The clinical assessment of the patient with acute localized abdominal pain, based on a careful history and examination together with examination of the urine, may be supplemented by laboratory and radiological investigations. A full blood count, plain X-ray of the abdomen, and estimation of the serum amylase in suspected pancreatitis may all be helpful, although, as mentioned above, the findings must be interpreted with caution. In suspected ruptured ectopic pregnancy, the urinary beta human chorionic gonadotrophin (beta hCG) is positive. Ultrasound of the pelvis may be helpful if a twisted ovarian cyst or some other pelvic pathology is suspected. Ultrasonography is also valuable in demonstrating gallstones in acute cholecystitis (Fig. A.2). An emergency intravenous urogram is indicated when a ureteric stone or some other renal pathology is suspected.

An electrocardiogram and appropriate cardiac enzyme estimations are performed if it is suspected that the upper abdominal pain is referred from a myocardial infarction, and a chest X-ray may demonstrate a basal pneumonia. Computed tomography is particularly useful in demonstrating the swollen and oedematous pancreas of acute pancreatitis (Fig. A.3). It must be stressed, however, that the clinical features take precedence over all other diagnostic aids.

Nothing can be simpler, or more difficult, than diagnosing a patient with the so-called ‘acute abdomen’. Particular difficulties will be encountered in infants (where history may be difficult and examining a screaming child most demanding), and in the elderly, where again it is often difficult to obtain an accurate history and where physical signs are often atypical.
Grossly obese individuals and pregnant women are two other categories where particular difficulties may be encountered.

When faced with a patient with severe abdominal pain, the main decision that must be taken, of course, is whether or not a laparotomy is indicated as a matter of urgency. If careful assessment still makes the decision difficult, repeated observations must be carried out over the next few hours to observe the trend of the particular case. This will nearly always enable a definite decision to be made on whether laparotomy or further conservative treatment is indicated.

**ABDOMINAL PULSATION**

Harold Ellis

A pulsatile swelling in the abdomen may be due to:
- A prominent aorta – normal or arteriosclerotic
- An abdominal aortic aneurysm
- Transmission of aortic pulsations through an abdominal mass
- A pulsatile, enlarged liver

**PROMINENT AORTA**

The pulsations of the normal aorta may be felt in perfectly normal but thin subjects along a line extending from the xiphoid to the bifurcation of the aorta at the level of the fourth lumbar vertebra. This is on a line joining the iliac crests, about 2 cm below and a little to the left of the umbilicus. In the arteriosclerotic and hypertensive subject, it may be difficult to decide whether or not the aorta is merely thickened and tortuous, or whether it is aneurysmal. If the two index fingers are placed parallel, one on either side of the aorta, the distance between the fingers can be measured. According to the size of the patient, a gap of 2–3 cm between the fingertips may be considered normal, but any measurement above this is suspicious of aneurysmal dilatation.

If in doubt, visualization of the aorta by means of ultrasound or computed tomography enables accurate measurement of the aorta to be made.

**ABDOMINAL AORTIC ANEURYSM**

There is no doubt that arteriosclerotic abdominal aneurysms are becoming more frequently encountered, as is the serious emergency of leakage or rupture of such an aneurysm. The majority of patients are aged more than 60 years, and the great majority are men. The aneurysm may be entirely symptomless or the patient may complain of epigastric or central abdominal discomfort that frequently radiates into the lumbar region. Patients themselves may actually detect the pulsating mass in the abdomen.

The pulsation may be visible in the upper abdomen, above the umbilicus, and – if large enough – may actually appear as a pulsating mass. On palpation, the aneurysm is a midline swelling that bulges over to the left side, away from the adjacent inferior vena cava. If the mass extends below the level of the umbilicus, it suggests implication of the iliac arteries. The characteristic physical sign is that the mass has an expansile pulsation. The index fingers are placed one either side of the mass, which enables the diameter to be assessed. If the diameter is more than 3 cm, this certainly suggests aneurysmal dilatation of the aorta; if the diameter is above 5 cm, the clinical diagnosis is all but certain. Typically, the fingers are pushed apart with each pulse, and not up and down. The latter sign suggests transmission of the pulsation (see section below).

Usually, the aneurysm is resonant to percussion due to overlying loops of intestine. However, an extremely large aneurysm will displace the bowel laterally to reach the anterior abdominal wall and will then give a dull percussion note. Auscultation may reveal bruits over the lower extremity of the aneurysm. This suggests turbulent flow of blood caused by relative stenosis at the aorto-iliac junctions.

Rectal examination may reveal a pulsatile mass when one or both of the internal iliac arteries are involved in the aneurysmal process.

Leakage or rupture of the aneurysm is an acute abdominal emergency. The patient presents with the features of massive blood loss (pale, sweating, clammy skin, a rapid pulse and low blood pressure) together with severe abdominal pain, lumbar pain and marked abdominal tenderness and guarding. Because of the low blood pressure and the associated peri-aneurysmal haematoma, as well as the overlying guarding, the aneurysm may be quite difficult to palpate and, unless sought carefully, is easy enough to miss.

The diagnosis of aortic aneurysm is often readily confirmed by means of a plain abdominal X-ray (Fig. A.4), which frequently delineates the aneurysm because of the associated calcification in its wall. Typically, the aneurysm is seen to bulge over to the left side of the abdomen. More accurately, an ultrasound or computed tomogram of the abdomen visualizes the aneurysm and enables its length and diameter to be measured accurately.
TRANSMISSION OF AORTIC PULSATIONS THROUGH AN ABDOMINAL MASS

A large intra-abdominal or retroperitoneal solid mass, pressing against the aorta, may exhibit transmitted aortic pulsation. Typical examples are a large carcinoma of the body of the stomach, a carcinoma or cyst of the pancreas, and a large ovarian cyst. Indeed, when the whole abdomen is filled by a cystic mass, it may be quite difficult to distinguish between such a mass and extensive ascites. Percussion, of course, is helpful since ascites gives dullness in the flanks as compared with the central dullness of a large intra-abdominal mass. The two index fingers, when placed on the mass, will perceive that the pulsation is transmitted directly forwards from the aorta and is not expansile, as would be found in an aneurysm.

PULSATILE LIVER

It is unlikely that an enlarged pulsatile liver will be mistaken for any other kind of pulsatile tumour. It occurs in cases of chronic failure of cardiac compensation, generally from mitral stenosis or tricuspid stenosis. There is associated cyanosis, oedema of the legs and ascites. It is not, however, every liver which seems to pulsate that really presents expansile pulsation. An impression of pulsation may be given by the movements transmitted directly to the liver by the hypertrophied right heart.

ABDOMINAL RIGIDITY

Harold Ellis

Rigidity of the abdomen is a sign of utmost importance, since in most cases it indicates serious intra-abdominal mischief requiring immediate operation. It is the expression of a state of tonic contraction in the muscles of the abdominal wall. The responsible stimulus may be in the brain or basal ganglia, or in the territory of the six lower dorsal nerves that supply the abdominal wall. The extent of the rigidity will depend on the number of nerves involved, and its degree on the nature and duration of the stimulus. The analysis in Table A.1 may be considered.

The patient should be examined lying on the back with the whole abdomen and lower thorax exposed, but with the shoulders and legs well covered. The room must be warm. The examiner, seated on a level with the patient, should first watch the abdomen to see whether it moves with respiration or not, and whether one part moves more than another; at the same time, he or she may observe other things that will help in the diagnosis, such as asymmetry of the two sides, local swelling, or the movement of coils of bowel.

Figure A.4 (a) Plain X-ray of the abdomen, showing a large calcified aortic aneurysm (arrowed). (b) Coronal computed tomography image of an infrarenal aortic aneurysm with a calcified wall (blue arrow) and intraluminal thrombus (red arrow). The arteries have been enhanced by an intravenous injection of contrast.
ABDOMINAL RIGIDITY

Table A.1 The extent of abdominal rigidity

<table>
<thead>
<tr>
<th>Site of stimulus</th>
<th>Causative agent</th>
<th>Characters of rigidity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cerebral cortex or basal ganglia</td>
<td>Nervousness, anticipation of pain, cold</td>
<td>Affects the whole abdominal wall; varies in intensity, can be abolished by appropriate means</td>
</tr>
<tr>
<td>Dorsal nerve trunks</td>
<td>Pleurisy, infections of the chest wall</td>
<td>Limited to one side of the abdomen; varies in extent and degree</td>
</tr>
<tr>
<td>Nerve endings in abdominal wall</td>
<td>Injury or infection of muscles</td>
<td>Limited to injured or infected segment</td>
</tr>
<tr>
<td>Nerve endings in peritoneum</td>
<td>Irritation by any intraperitoneal foreign substance: infection, chemical irritant, or blood</td>
<td>Degree varies with nature of irritant and suddenness with which stimulus has arrived. Extent corresponds to area of peritoneum involved. Both degree and extent remain approximately constant during the period of examination</td>
</tr>
</tbody>
</table>

While watching – and later when examining – the examiner should engage the patient in conversation, encouraging him to talk in order to allay nervousness and to remove any part of the rigidity that is due to a voluntary contraction. Some nervous patients – especially if the room is cold – hold their abdomens intensely rigid, and can be induced to relax only after gentle persuasion; a request to take a few deep breaths, or to draw the knees up and keep the mouth open, will often help.

During this preliminary examination, one (well-warmed) hand may be laid gently on the abdomen and passed over its surface with a light touch that cannot possibly hurt; this manoeuvre will help to allay the patient’s anxiety still further and give the examiner an idea of the extent, intensity and constancy of the rigidity to be investigated later in more detail.

For a more exact examination, the observer should sit at the patient’s side facing their head, and place both hands on the abdomen, examining comparable areas of both sides, simultaneously, and taking in turn the epigastrum, right and left hypochondria, umbilical region, both flanks as far back as the erector spinae (as the rigidity of a retrocaecal appendix may only affect the posterior part of the abdominal wall), the hypogastrium and both iliac fossae. First, the whole hand should be applied with light pressure; next, the fingers held flat should be pressed more firmly to estimate the extent of the rigidity and to discover deep tenderness; last, a detailed examination may be made in suspected areas with the firm pressure of one or two fingers. Evidence is not complete without percussion and auscultation. A rectal examination is indispensable.

After a leisurely examination with warm hands in a warm room, during which the physician has also been able to sum up the patient, their temperament, and whether they are really ill or not, the rigidity of anxiety or cold will have been dispelled or recognized.

The abdominal rigidity due to a lesion in the chest or chest wall usually involves a wide area limited to one side – a distribution most unusual with intra-abdominal mischief, which, if it has spread widely but not everywhere, tends to be limited to the upper or lower half. The extent and degree of rigidity in chest affections also vary widely during examination. Other things such as a flushed face, rapid respiration, movement of the alae nasi, or a temperature of more than 39 °C (102 °F) may suggest that the lesion is not abdominal, and a friction rub may be felt or heard in the chest.

Auscultation and rectal examination dispel any remaining doubts, as in chest conditions peristaltic sounds remain normal and there is no tenderness in Douglas’s pouch.

Examination of the blood may show a high leucocytosis (up to 30 000 or 40 000 per mm³), whereas in peritonitis the count is seldom over 12 000 per mm³. Chest X-rays (including a lateral film) will demonstrate the intrathoracic lesion.

Injuries of the abdominal wall, and particularly those caused by run-over accidents, lead to very marked rigidity of the injured segment. Here, the rigidity is not necessary to establish a diagnosis, as the injury is already known, but its degree and extent should be carefully noted. There must always be a doubt as to whether the abdominal viscera are damaged as well as the walls, and this point can only be settled by careful observation. The patient is put to bed and kept warm, the pulse is charted every 15 minutes, and the abdomen is re-examined from time to time. In the case of a mere contusion, the collapse will soon disappear, the abdomen will become less rigid, and the pulse rate will fall. If the contents of a hollow viscus have escaped, rigidity will extend beyond the area of the damaged muscles, and the signs of peritonitis will develop rapidly. An X-ray of the abdomen, in the erect position, will demonstrate free gas beneath the diaphragm (see Fig. A.1). If there is internal bleeding.
The development of a case. The board-like abdominal wall

The degree of muscle contraction also alters during the

blood, still less. Bacterial invasion of the peritoneum

The extent of the rigidity usually corresponds to the

increased resistance when the hands are pressed on

there is a slighter resistance when the muscles contract over a wide

increases as infection supervenes.

perforation, pancreatitis or the bursting of a large

amylase in acute pancreatitis leads to less rigidity, and

The influence of natural subdivisions in guiding

of a perforation is considerably softer after 3–4 hours

When gallstone colic is followed by rigidity of the

The most common and the most important cause of

gives colic referred to the umbilicus but no

the presence of rigidity therefore

irritant, the rapidity with which the peritoneum

a local inflammatory focus such as

of bowel. In appendicitis, rigidity denotes that infection

locality, to make a diagnosis.

Local peritonitis starts around some site of infection,

Thus, localized rigidity is found over any inflamed

Excessive and the muscles contract over a wide area

that there is also a local inflammatory focus such as

organ, and as the infection and the guarding spread,

and as it spreads it is guided by certain peritoneal

thus, it stretches if the hands are pressed on

of the appendix, although this exaggerated response

infected fluid is suddenly

of the appendix lining the abdominal cavity is in contact with something

of peritonitis until it can be excluded. Actually, rigidity means

no more than that the parietal peritoneum lining

announces a change in the coelomic cavity that is

is blocking the cystic duct, but also that the wall of

Intestinal obstruction

there is no more than that a stone

of mechanical origin (such as that due to a band or

and its smooth surfaces that are its normal

parietal peritoneum lining the abdominal cavity is in contact with something differing from the smooth surfaces that are its normal environment. The presence of rigidity therefore

When gallstone colic follows by rigidity of the
differs from the smooth surfaces that are its normal

When gallstone colic follows by rigidity of the

tends to pass down between the ascending colon and

of peritonitis, and it is a safe

When gallstone colic follows by rigidity of the

tends to pass down between the ascending colon and

Thus, localized rigidity is found over any inflamed

locality, to make a diagnosis.

Local peritonitis starts around some site of infection,

Thus, localized rigidity is found over any inflamed

and its smooth surfaces that are its normal

when the peritoneum has recovered from the shock

of a perforation is considerably softer after 3–4 hours

of a perforation is considerably softer after 3–4 hours

pancreatitis or peritonitis until it can be excluded. Actually, rigidity means

no more than that the parietal peritoneum lining

Intestinal obstruction

that there is also a local inflammatory focus such as

Intestinal obstruction

that there is also a local inflammatory focus such as

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Intestinal obstruction

where the abdomen is suddenly flooded with gastric

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ABDOMINAL RIGIDITY

rectal examination. Percussion may reveal the outline of some dilated hollow organ, such as the caecum; it may disclose free gas that has escaped from a perforation as a shifting circle of resonance or a tympanic note where liver dullness should be; it may map out an abnormal area of dullness where there is an abscess or a collection of blood; or it may indicate free fluid in the peritoneum. Auscultation is even more important, as peristalsis ceases with peritonitis: in a normal abdomen, peristaltic sounds can be heard every 4–10 seconds; in obstruction, they are increased in loudness, pitch and frequency; but in peritonitis, there is complete silence. Rectal examination nearly always reveals tenderness when there is intra-abdominal infection, even if it is distant and localized.

Other signs must be mentioned: the patient lies still, sometimes with the knees drawn up, and resists interference. The abdomen gradually becomes distended, tense and tympanitic. The tongue is brown and dry. Vomiting is to be expected at the onset of any abdominal catastrophe, but it usually ceases, except in intestinal obstruction. With advancing peritonitis, it reappears, and the vomit becomes first bile-stained, later brownish and faecal-smelling, and is allowed to dribble from the corner of the mouth in contrast to the projectile vomiting of obstruction. There may be diarrhoea at first, but absolute constipation soon succeeds it. The temperature tends to fall; the pulse is small and rapid, rising progressively. In late stages, the sunken cheeks, wide eyes and anxious expression of the patient form a characteristic feature – the Hippocratic facies. These signs are indications of a peritonitis discovered too late, and are the heralds of approaching death. Abdominal rigidity, abdominal silence, rectal tenderness and a rising pulse are a tetrad that calls for immediate definitive treatment.

A more detailed diagnosis is usually possible when the history and other signs are taken together, but a consideration of all the alternatives is out of the question in this section. Abdominal paracentesis with a fine needle may clinch the presence of pus, blood or urine in the peritoneal cavity, but a false-negative tap may delay rather than aid diagnosis. A list of the more common conditions associated with rigidity may, however, help the inquiry:

- Stomach or duodenum
  - Perforation of peptic ulcer
- Gallbladder
  - Acute cholecystitis
  - Rupture of the gallbladder
- Pancreas
  - Acute pancreatitis
- Small intestine
  - Strangulation of a loop
  - Traumatic perforation
  - Mesenteric vascular thrombosis or embolism
  - Meckel's diverticulitis
  - Acute ileitis
- Large intestine
  - Appendicitis
  - Volvulus
  - Diverticulitis with perforation
- Peritoneum
  - Acute blood-borne peritonitis
    - Streptococcal
    - Pneumococcal
    - Gonococcal
- Female generative organs
  - Twisted ovarian cyst
  - Ruptured ectopic pregnancy
  - Acute salpingitis
  - Torsion or red degeneration of a fibroid
  - Perforation of the uterus or posterior fornix of vagina in attempted abortion
- Spleen and/or liver
  - Traumatic rupture
- Aorta
  - Ruptured aneurysm

Perforation of a peptic ulcer is characterized by the most sudden onset, the worst agony and the most extreme abdominal rigidity that the physician is ever likely to see. Radiation of pain to the right shoulder tip (referred pain from diaphragmatic irritation) may be experienced. Immediately afterwards, the patient is motionless and speechless, in a state of obvious collapse. A few hours later, pain, rigidity and shock have all diminished, and only the dramatic history of sudden onset and persistent abdominal and rectal tenderness may remain to indicate the seriousness of the condition.

Acute pancreatitis is seldom accompanied by the severe pain described in textbooks, or indeed by pain as bad as that of gallstone colic. The abdominal rigidity is more marked in the upper abdomen but is not profound. On the other hand, the patient shows a degree of toxaemia out of all proportion to the physical signs in the abdomen. The diagnosis is confirmed by a considerable rise in the serum amylase (see p. 4).

A ruptured ectopic pregnancy may simulate a lower abdominal peritonitis, but the signs of bleeding predominate and rigidity is not well marked. If the patient is a woman of child-bearing age who is known to have missed a period, the onset of abdominal pain and pallor suggest the diagnosis. Extravasated blood
will be felt in the pelvis, together with acute tenderness on vaginal and rectal examinations. The diagnosis is confirmed by the urinary pregnancy test (positive beta human chorionic gonadotrophin (hCG)) and, if necessary, by pelvic ultrasonography.

Blue discoloration of the skin around the umbilicus – Cullen’s sign – may be associated with rigidity. This discoloration is due to extravasated blood coming forwards from the retroperitoneal space. The sign is seen in ruptured kidney, leaking abdominal aneurysm and acute pancreatitis. Occasionally, it is seen in ruptured ectopic pregnancy, when the blood gains entry to the subperitoneal space through the broad ligament. Although pancreatitis may produce this sign, it is more common to see a green discolouration in the loins (Grey Turner’s sign).

**ABDOMINAL SWELLINGS**

Harold Ellis

*(See also VEINS, VARICOSE ABDOMINAL, p. 735.)*

These may be acute or chronic, general or local, and caused by abdominal accumulations that are gaseous, liquid or solid. They may arise in the abdominal cavity itself or in the abdominal wall.

**SWELLINGS IN THE ABDOMINAL WALL**

Swellings situated in the abdominal wall itself can be recognized by their superficial position, by their adherence to the skin, subcutaneous fascia or muscles, or by their failure to follow the movements of the viscera immediately underlying the abdominal wall (Fig. A.5). It may be impossible to differentiate, for obvious reasons, an intra-abdominal mass that has become attached to the abdominal parietes, either as an inflammatory or malignant process. A simple test that should be applied to all abdominal masses is to ask patients to raise either their legs or their shoulders from the couch. This procedure tightens the abdominal muscles; if the lump is intraperitoneal, it disappears, but if it is situated in the abdominal wall itself it persists.

**Inflammatory swelling** of the abdominal wall most commonly complicates a laparotomy incision, and the diagnosis is obvious. A superficial cellulitis may complicate infection of a small abrasion or hair follicle infection. Inflammation of the abdominal wall may be secondary to an extension of an intraperitoneal abscess, particularly an appendix abscess in the right iliac fossa, or, on the left side, a paracolic abscess in relation to diverticular disease of the sigmoid colon or to perforation of a carcinoma of the large bowel.

**Inflammatory swelling of the umbilicus** in newborn infants is rare, except in primitive communities where the cord is not divided with the niceties of modern aseptic practice. Suppuration at the umbilicus in adults is not uncommon if the navel is deep and narrow. A tender haematoma in the lower abdomen may result from rupture of the rectus abdominis muscle, or tearing of the inferior epigastric artery, which may occur as the result of a violent cough.

**Tumours** of the abdominal wall are usually subcutaneous lipomas. These may be multiple and may be a feature of Dercum’s disease (adiposa dolorosa). Lipomas should be carefully differentiated from irreducible umbilical or epigastric hernias containing omentum.

A desmoid tumour may arise in the lower part of the abdominal wall, and malignant fibrosarcomas or melanomas may also occasionally be encountered. A neoplastic deposit may sometimes be palpated at the umbilicus and represents a transcoelomic seeding, usually from a carcinoma of the stomach or large bowel.

**GENERAL ABDOMINAL SWELLINGS**

Every medical student knows the mnemonic of the five causes of gross generalized swelling of the abdomen: Fat, Fluid, Flatus, Faeces and Fetus.

In *obesity*, the abdomen may swell either in consequence of the deposit of fat in the abdominal wall itself, or as the result of adipose tissue in the mesentery, the omentum and the extraperitoneal layer. In very obese persons, it is rarely possible to diagnose the exact nature of an intra-abdominal mass by the usual clinical methods. Indeed, tumours of quite remarkable size – including the full-term fetus – may remain occult to even the most careful examiner.
Distension of the intestines with gas occurs in intestinal obstruction and is particularly marked in cases of volvulus of the sigmoid colon, chronic large-bowel obstruction and megacolon. It also occurs in an adynamic ileus. The whole of the abdomen, or in special cases some part of it, is distended and gives on percussion a highly resonant or tympanitic note. The outlines of the gas-distended viscera are often visible; loops of dilated small bowel, one above the other, may produce a characteristic ‘ladder pattern’. The increased size of the inflated intestine may produce displacement of the other viscera; the dome of the diaphragm is pushed up into the chest, shifting the apex beat of the heart upwards. The liver is similarly displaced. The distended stomach may occasionally be gross enough all but to fill the abdomen in very advanced cases of pyloric stenosis and in acute gastric dilatation.

The causes producing an accumulation of liquid in the peritoneal cavity can be listed as:

- Congestive cardiac failure
- Cirrhosis
- Nephrotic syndrome
- Carcinomatosis peritonei
- Tuberculous peritonitis

In severe cases of chronic constipation, abdominal distension may result from the accumulation of faeces in the large intestine, particularly where megacolon exists. The scybala may be felt, usually soft and plastic in the region of the ascending colon, and hard and nodular in the descending and sigmoid colon. Rectal examination often reveals an enormous accumulation of faeces. In some cases of tuberculous peritonitis, semi-solid inflammatory masses may bring about a general swelling of the abdomen. General swelling of the abdomen may occur in malignant disease involving the peritoneum due to the growth of numerous secondary nodules in addition to a concomitant ascites. *Pseudomyxoma peritonei* may follow rupture of a pseudomucinous cystadenoma of the ovary or of a mucocoele of the appendix. The whole abdominal cavity becomes distended with gelatinous material.

**LOCAL INTRA-ABDOMINAL SWELLINGS**

These may be due to some general cause, or to a mass arising in a specific viscus.

**Swellings due to general causes**

Causes that ordinarily produce general swelling of the abdomen may sometimes give rise to only a local swelling. Thus, with encysted ascites left after an acute diffuse peritonitis or accompanying tuberculous peritonitis, an accumulation of fluid bounded by adhesions between the adjacent viscera may be found in any part of the peritoneal cavity, most often in the flanks or pelvis. A reliable history may be a clue to the nature of such a mass, although its cause may not be revealed until a laparotomy has been performed.

Abdominal swellings may occur in tuberculous peritonitis resulting from the rolled-up, matted and infiltrated omentum, doughy masses of adherent intestine, or enlarged tuberculous mesenteric lymph nodes. The amount of ascites in such cases varies considerably from a gross degree to almost complete absence (the obliterator form). Discovery of a tuberculous focus elsewhere in the body is support for the diagnosis.

*Hydatid cysts may occur in any part of the abdominal cavity. They are usually single. The liver – particularly the right lobe – is the most common situation, and more rarely the spleen, omentum, mesentery or peritoneum. The cyst grows slowly and is spherical except in so far as it is moulded by the pressure of adjacent structures. It contains a clear fluid in which may be found hooklets, scolices and secondary or daughter cysts detached from the walls of the parent cyst.

 Unless large enough to cause mechanical pressure, the single hydatid cyst gives rise to little pain, or indeed to any complaint of any kind. It may produce a smooth, rounded, tense bulging of the overlying abdominal wall. It is dull on percussion, and it may yield a ‘hydatid thrill’, as may any other cyst; this thrill is the vibratory sensation experienced by the rest of the hand when, with the whole hand laid flat over the tumour, a central finger is percussed. Occasionally, there may be pain and fever due to inflammation within these cysts, and rupture into the peritoneal cavity may cause a severe anaphylactic reaction. Rupture of a hydatid cyst of the liver into a bile duct may cause jaundice due to biliary obstruction by daughter cysts. Hydatid disease is rare except in countries where the inhabitants live in close association with dogs that are the hosts of *Taenia echinococcus* (Australasia, South America, Greece, Cyprus and, in the British Isles, North Wales). About one-quarter of patients demonstrate eosinophilia. A complement fixation test gives a high degree of accuracy. X-rays of the abdomen may reveal calcification of the cyst wall in long-standing cases.

Any part of the abdomen may swell from the formation of an abscess. A subphrenic abscess following a general peritonitis is occasionally large enough to produce an upper abdominal swelling. The patient is usually seriously ill with a swinging fever, rapid pulse, leucocytosis and all the general...
manifestations of toxaemia. However, in this antibiotic era, an increasing number of examples are being seen of a more insidious and chronic progress of the disease, with the onset delayed weeks or even many months after the initial peritoneal infection.

X-ray examination, together with screening of the diaphragm, is extremely useful, and at least 90 per cent of patients with subphrenic infection have some abnormality on this investigation. On the affected side, the diaphragm is raised and its sharp definition is lost. Its mobility on screening is diminished or absent. There is frequently a pleural effusion, collapse of the lung base or evidence of pneumonitis. About 25 per cent of patients have gas below the diaphragm, frequently associated with a fluid level. This gas is usually derived from a perforated abdominal viscus, but it is occasionally formed by gas-producing organisms. On the left side, gas under the diaphragm may be confused with the gastric bubble. An important differential feature is that the gas shadow of the stomach rarely reaches the lateral abdominal wall; however, if there is doubt, a mouthful of barium is given in order to demarcate the stomach. Ultrasonography and computed tomography usually clinch the diagnosis.

Pus may localize in either the right or left paracolic gutter or iliac fossa. On the right side, this commonly follows a ruptured appendix, or occasionally a perforated duodenal ulcer. On the left, a perforation of an inflamed diverticulum or carcinoma of the sigmoid colon is the usual cause. A large pelvic abscess frequently extends above the pubis or into one or other iliac fossa from the pelvis and can be palpated abdominally as well as on pelvic or rectal examination. About 75 per cent result from gangrenous appendicitis, and the remainder follow gynaecological infections, pelvic surgery or any general peritonitis.

Regional diagnosis of local abdominal swellings
For clinical purposes, the abdomen may be subdivided into nine regions by two vertical lines drawn upwards from the mid-inguinal point midway between the anterior superior iliac spine and the symphysis pubis, and by two horizontal lines, the upper one passing through the lowest points of the tenth ribs (the subcostal line), the other drawn at the highest points of the iliac crests – the supra-cristal plane (Fig. A.6).

The three median areas thus mapped out are named, from above downwards, the epigastric, umbilical and hypogastric (or suprapubic) regions; the six lateral areas are, from above downwards, the right and left hypochondriac, lumbar and iliac regions.

Figure A.6 The regions of the abdomen. Identification numerals are listed in Box A.1.

The viscera, or portions of viscera, commonly situated in the areas thus demarcated are listed in Box A.1. The abdominal swellings that may be felt in and about these nine regions, excluding the tumours situated in the abdominal wall itself that have already been described, are as follows.

**Right hypochondriac region**
Most tumours in this area are connected with the liver or gallbladder, and their differential diagnosis is discussed under LIVER, ENLARGEMENT OF (p. 347) and GALLBLADDER, PALPABLE (p. 213).

An easily made mistake is to regard the firm and rounded swelling produced by the upper segment of the right rectus abdominis muscle, especially in a well-developed subject, as a tumour of the liver or gallbladder. In such cases, the characteristic dull note of the liver on percussion over the lower right chest ceases at the costal margin.

Tumours in connection with the hepatic flexure of the colon, scybalous collections in the hepatic flexure region, or the head of an intussusception may present as masses in this area.
Epigastric region

Enlargement of the liver may be felt in this area, and indeed it is common to feel the normal liver in this region, especially in infants and in adults with an acute costal angle. The dilated stomach produced by pyloric stenosis in either children or adults may present as a visible swelling demonstrating waves of peristalsis traveling from left to right (Fig. A.7). A succession splash is usually elicited. Tumours of the stomach, apart from malignant growth, are rare. A hundred years ago, a hair ball or trichobezoar was frequently encountered as an epigastric mass in hysterical girls who chewed and swallowed their hair, which then formed an exact mould of the stomach. Hair balls are only rarely encountered these days, and modern textbooks hardly mention them; however, as fashions and hair styles change, they may reappear on the clinical scene (Fig. A.8). Other foreign bodies are sometimes ingested by those with learning difficulties and form a palpable mass. In congenital pyloric stenosis, a tumour the size of a small marble is palpable at the right border of the right rectus.

### Box A.1 The normal contents of the abdominal regions

<table>
<thead>
<tr>
<th>Region</th>
<th>Contents</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Right hypochondriac region</td>
<td>Liver, Gallbladder, Hepatic flexure of colon, Right kidney, Right suprarenal gland</td>
</tr>
<tr>
<td>2. Epigastric region</td>
<td>Liver, Stomach and pylorus, Transverse colon, Omentum, Pancreas, Duodenum, Kidneys, Suprarenal glands, Aorta, Lymph nodes</td>
</tr>
<tr>
<td>3. Left hypochondriac region</td>
<td>Liver, Stomach, Splenic flexure of colon, Spleen, Tail of pancreas, Left kidney, Left suprarenal gland</td>
</tr>
<tr>
<td>4. Right lumbar region</td>
<td>Riedel's lobe of the liver, Ascending colon</td>
</tr>
<tr>
<td>5. Umbilical region</td>
<td>Stomach, Duodenum, Transverse colon, Omentum, Urachus, Small intestine, Aorta, Lymph nodes</td>
</tr>
<tr>
<td>6. Left lumbar region</td>
<td>Descending colon, Small intestine, Left kidney</td>
</tr>
<tr>
<td>7. Right iliac fossa</td>
<td>Caecum, Vermiform appendix, Lymph nodes</td>
</tr>
<tr>
<td>8. Hypogastric region</td>
<td>Small intestine, Sigmoid flexure, Distended bladder, Urachus, Enlarged uterus and adnexa</td>
</tr>
<tr>
<td>9. Left iliac fossa</td>
<td>Sigmoid flexure, Lymph nodes</td>
</tr>
</tbody>
</table>

Figure A.7 Grossly distended upper abdomen, which showed visible peristalsis from left to right in a patient with pyloric stenosis due to a chronic duodenal ulcer.

Figure A.8 Gastric hair ball (trichobezoar). This formed a large mobile epigastric mass in a young woman with long hair. (a) The mass being removed at gastrotomy. (b) The removed specimen.

The transverse colon usually passes across the upper part of the umbilical area, and may be palpated when it is the site of a carcinoma, when it is impacted.
with faeces or when it is distended by a large-bowel obstruction placed distal to it.

Swellings in connection with the *omentum* may be due to tuberculous peritonitis or, more commonly, due to infiltration with secondary malignant deposits.

Swellings arising from the *pancreas* push forward from the depths of the abdominal cavity towards the epigastric and the upper part of the umbilical areas, and present themselves as vaguely palpable deeply seated masses. They have the stomach, or the stomach and colon, in front of them and are fixed to the posterior abdominal wall, thus moving only a little on respiration. They may transmit a non-expansile pulsation from the subjacent aorta. Unless extremely large, such swellings are resonant on percussion, due to the overlying air-filled gut. A pancreatic swelling may be carcinomatous, in which case wasting, anaemia and jaundice are likely to be observed. There may be clay-coloured stools and dark urine, and it is important to note that the onset of jaundice is frequently preceded by deeply placed abdominal pain, or pain in the back. Glycosuria of recent origin in an elderly patient also raises suspicion of a pancreatic carcinoma. In about half the patients with jaundice due to carcinomatous obstruction, the gallbladder is palpably distended (Courvoisier’s law) (see Fig. G.4). Occasionally, the mass may result from chronic pancreatitis; the swollen pancreas of acute pancreatitis has only exceptionally been palpated before laparotomy.

Pancreatic cysts are the pancreatic swellings that are most commonly palpable. Only 20 per cent are true cysts; these are either single or multiple retention cysts that usually result from chronic pancreatitis, neoplastic cysts (cystadenoma and cystadeno-carcinoma) and the rare congenital polycystic disease of the pancreas and hydatid cyst of the pancreas. Far more often, the cysts are not in the pancreas itself but comprise a collection of fluid sealed off in the lesser sac due to closure of the foramen of Winslow (pseudocyst of the pancreas). This may occur after trauma to the pancreas, following acute pancreatitis or, much less commonly, resulting from perforation of a posterior gastric ulcer. They may reach an enormous size and fill the whole upper part of the abdomen.

*Retroperitoneal cysts* are rare. The majority arise from remnants of the mesonephric (Wolffian) duct and occur in adult women. Others are teratomatous, lymphangiomatous or dermoid.

Retroperitoneal tumours (apart from those arising in the pancreas, suprarenal gland or kidney) originate in the mesenchymal tissues, the sympathetic chain and the para-aortic lymph nodes.

Swellings in connection with the *duodenum* are exceedingly rare. They may result from an inflammatory mass developing around a penetrating duodenal ulcer, or be due to a duodenal malignant tumour, but the latter is a pathological curiosity. Those in connection with the *kidneys* and *suprarenal glands* are found in the epigastrium only if very large. Their diagnosis is considered below.

Enlargement of the *spleen* may bring its anterior edge into the epigastric area; a splenic swelling always lies in contact with the anterior wall of the abdomen (see SPLENOMEGALY, p. 635).

Lymph nodes, which are numerous in the para-aortic retroperitoneal tissues and in the mesentery, may become palpable in reticulosus, tuberculous peritonitis, or malignant disease as nodulated chains or masses.

**Left hypochondriac region**

An abnormal lobe or a tumour in the left lobe of the *liver* may appear as a superficial tumour in this area. Much of the *stomach* normally lies in the left hypochondrium; the diagnosis of gastric swelling has been considered above, and a gastric tumour is commonly felt in this region. On physical signs alone, it must be differentiated from a swelling of the adjoining *spleen*. A barium-meal X-ray examination, ultrasound or computed tomography (CT) scan help considerably in differentiating between a gastric and a splenic swelling.

The diagnosis of a tumour of the splenic flexure of the *colon*, whether scabby or malignant, is arrived at in the same way as in the case of a tumour of the hepatic flexure or transverse colon (see ‘Right hypochondriac region’ and ‘Epigastric region’, above).

The diagnosis of various causes of enlargement of the *spleen* is discussed under SPLENOMEGALY (p. 635). The distinguishing features are that the spleen comes down from under the left costal margin in direct contact with the anterior abdominal wall (and is therefore dull on percussion), descends on inspiration and has a smooth surface, and a notch may be palpable on its inner margin. A splenic swelling may be identified on a plain X-ray of the abdomen and differentiated from a renal mass by means of pyelography. A barium meal examination may show displacement and indentation of the adjacent stomach. Ultrasound or CT scan will clinch the diagnosis.

Tumours of the *pancreas* may project into the left hypochondrium, as may retroperitoneal tumours and cysts (see ‘Epigastric region’, above).

Tumours of the left *kidney* and *suprarenal gland* have the stomach and colon in front of them and therefore,
ABDOMINAL SWELLINGS

unless extremely large, are resonant on percussion. Since they arise in the loin, these masses can usually be ballotted by bimanual palpation.

**Right lumbar region**

Occasionally, a congenital projection of the liver, known as Riedel’s lobe, may appear as a superficial tumour continuous with the liver above it in this zone. It may be mistaken for a dilated gallbladder.

The ascending colon may be palpable due to contained faecal masses, owing to thickening as a result of long-standing colitis, Crohn’s disease or hyperplastic tuberculosis, or due to malignant disease.

The ascending colon can be felt in acute or chronic ileocaecal and ileocolic intussusception as a sausage-shaped tumour, at first situated in the right flank, then moving across the abdomen above the umbilicus and finally down the left flank into the pelvis. The vast majority of these cases occur in infants or young children, commonly aged between 3 and 12 months. Boys are affected twice as often as girls. The history is of paroxysms of abdominal colic typified by screaming and pallor. There is vomiting and usually the passage of blood and mucus per rectum, giving the characteristic ‘redcurrant jelly stool’. A rectal examination nearly always reveals this typical feature, and rarely the tip of the intussusception can be felt. In infants, there is usually no obvious cause, but the mesenteric lymph nodes in these cases are invariably enlarged.

In adults, a polyp, carcinoma or an inverted Meckel’s diverticulum may form the apex of the intussusception. Tumours in connection with the right kidney and suprarenal gland usually appear deep down in this region, having the ascending colon and small intestine in front of them. They can be lifted forwards en masse from behind by a hand placed at the back of the loin and thus palpated bimanually. For their diagnosis, see KIDNEY, PALPABLE (p. 334). The lower pole of the right kidney can be felt in some cases of persons on deep abdominal palpation, especially in thin females. When abnormally low and mobile, the whole of the otherwise normal kidney may be palpable. Its shape and consistency are characteristic. Renal swellings move on respiration; unless very large, are resonant on percussion due to the anteriorly related gut. However, Riedel’s lobe of the liver, an enlarged gallbladder, masses in the ascending colon and secondary deposits in the omentum have all been mistaken for it, although they are more superficially placed and lie in contact with the anterior abdominal wall. Other wandering masses, for example those arising from the ovary, Fallopian tube and mesentery, as well as hydatid cysts, are all liable to the same error of identification.

Imaging by means of ultrasound or CT scanning is invaluable in assistance with the differential diagnosis.

**Umbilical region**

The grossly dilated stomach resulting from long-standing pyloric obstruction may occupy the umbilical region; indeed, it may descend below it down into the pelvis. Tumours in connection with the transverse colon have been considered in 'Epigastric region' and 'Right lumbar region', above.

Tumours in connection with the omentum are common in this region; those arising from the small intestine are much rarer, although the thickened small bowel in Crohn’s disease may form a palpable mass.

Swellings arising from the kidneys, suprarenal glands, pancreas, retroperitoneal tissues, para-aortic nodes and mesentery may all present themselves in the deeper parts of the umbilical region, usually as more or less fixed masses arising from or connected with the posterior wall of the abdomen.

The aorta bifurcates 1 cm below and to the left of the umbilicus in the supracristal plane, as shown in Figure A.6, above (at the level of the fourth lumbar vertebra). In thin patients, pulsation of the normal aorta can often be felt and indeed seen in this region, and may lead to the incorrect diagnosis of an abdominal aneurysm. Careful examination, however, will show that this pulsation is no more than a throbbing, an up-and-down movement, and is not laterally expansile. Aneurysm of the abdominal aorta forms an expansile mass situated above the umbilicus itself, and it may be accompanied by pain in the back from erosion of the bodies of the lumbar vertebrae. Often, X-rays of the abdomen in such cases will reveal calcification in the aneurysmal wall. Ultrasound and CT enable accurate delineation of the size and extent of the aneurysm. These methods are also valuable in the visualization of the other retroperitoneal masses enumerated above.

**Left lumbar region**

An enlarged spleen (see ‘Left hypochondriac region’, above) may protrude into this area. It forms a firm mass that is in contact with the abdominal wall, and its dullness to percussion continues with its thoracic dullness, which extends back up into the axilla along the line of the ninth or tenth ribs. Tumours in connection with the right kidney, the right suprarenal gland and the descending colon give rise to features similar to those considered in ‘Left hypochondriac region’, above.

**Right iliac fossa**

An inflammatory mass in this region is most commonly associated with an appendix abscess.
Less commonly, there may be a *paracaeal abscess* in relation to a perforated carcinoma of the caecum, or a solitary caecal benign ulcer. A *pyosalpinx* may result from salpingitis and, rarely, inflammatory swellings may arise in connection with suppurating *iliac lymph nodes* or a *psosas abscess*.

An important differential diagnosis is between an appendix mass and a carcinoma of the caecum. In the former, there is usually a preceding episode of an acute abdominal pain, typical of appendicitis, with fever and leucocytosis. The inflammatory mass subsides progressively over 2–3 weeks, and the occult blood test in the stools is negative. A carcinoma of the caecum may be suspected if there is a preceding history of bowel disturbance in a middle-aged or elderly patient, if the mass fails to resolve rapidly and if the occult blood test in the stools is repeatedly positive. If there is any clinical doubt, a barium enema X-ray examination should be carried out and, if necessary, resort made to laparotomy.

It is not at all rare for a soft ‘squelchy’ caecum to be palpable in a perfectly normal thin female subject. Occasionally, a grossly distended *gallbladder* may project down as far as the right iliac fossa, and a low-lying *kidney* may form a palpable mass in this region. Rarely, an ectopic kidney may be felt in one or other iliac fossa and, these days, a transplanted kidney may be palpated at this site. An *ovarian tumour* or cyst or a pedunculated *fibroid* of the uterus may project into this area.

**Hypogastric region**

The most common mass to be felt in this region, after the pregnant uterus, is the distended *bladder*. This may reach as high as, or slightly above, the umbilicus (see Fig. U.29). Not uncommonly, this midline structure tilts over to one or the other side. A distended bladder has been tapped as ascites, operated upon as an ovarian cyst or a fibroid, or mistaken for the pregnant uterus. No diagnostic opinion should be advanced, and no operative procedure undertaken respecting a tumour in this situation, until the bladder has been emptied, either by voluntary micturition or by the passing of a catheter.

Abdominal swellings arising from the *uterus, ovaries, Fallopian tubes* and *uterine ligaments* may all rise up out of the pelvis and present themselves as swellings in this region; as they grow larger, they may be spread into any part of the abdomen. While they remain comparatively small and are manifestedly connected with some intrapelvic organ, their origin is not difficult to determine (see PELVIS, SWELLING IN, p. 498). However, when they have extended into the abdomen or have acquired a long pedicle, or have become fixed by adhesions to some distant part of the abdominal wall or to some other viscus, these pelvic tumours may give rise to signs and symptoms that bear no relation to pelvic disease. In such cases, they may only be correctly diagnosed at laparotomy. The discerning clinician will always remember the possibility of pregnancy in every female patient between the menarche and menopause. The diagnosis is confirmed by the urinary pregnancy test (positive beta human chorionic gonadotrophin (hCG)) and, if necessary, by pelvic ultrasonography.

Tumours of ileal Crohn’s disease arising in the *small intestine* may be felt in the hypogastric area.

The *urachus* is a fibrous cord running in the middle line in front of the peritoneum from the fundus of the bladder to the umbilicus. Occasionally, it becomes the seat of cyst formation, more often in women than in men. The urachal cyst is a rounded tumour lying between the umbilicus and the pubic symphysis, which occasionally becomes infected.

**Left iliac fossa**

The *pelvic colon* can often be felt in normal subjects as a tube-like cord, either when empty and in spasm, or else when distended with faecal masses. The region is a common site for carcinoma of the colon, and there are usually symptoms of chronic intestinal obstruction, or bowel disturbance with the passage of blood and mucus in the stools. It is clinically impossible to differentiate between such a mass and that associated with diverticular disease of the sigmoid colon. Similarly, a paracolic abscess in this region may equally well be associated with suppuration of an inflamed colonic diverticulum or a perforating carcinoma. Rarely, such an abscess may be due to perforation of the tip of a long *appendix* passing over the left iliac fossa, or as an extreme rarity due to local perforation of a left-sided appendix in transposition of the viscera. The diagnosis of this would be suggested by finding the cardiac apex beat to lie on the *right* side.

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**ALCOHOL**

*Andrew Hodgkiss*

While some patients readily declare alcohol misuse, many do not. There are a number of common presentations that oblige the doctor to enquire carefully about the possibility of alcohol misuse. These are most readily grouped into *medical* (e.g. falls, fits, head injuries, haematemesis or jaundice), *psychiatric* (e.g. panic attacks, amnesic black-outs, confusional states and deliberate self-harm) and *social* (e.g. road traffic accidents, as the victim or perpetrator...
of violent crime, domestic violence or rough-sleeping). These may be the current presenting complaint or prominent in the past history. It is estimated that up to 20 per cent of UK medical admissions are for conditions caused by alcohol misuse, yet too few medical admissions have their drinking habits adequately assessed.

Assessment of alcohol misuse has three aims: to quantify use; to catalogue any alcohol-related problems the patient has; and to detect alcohol dependence syndrome if present.

Quantifying use by direct questioning is not always doomed to fail. The aim is to establish how many units the individual consumes in a typical week (or in a ‘heavy session’ if the pattern is binge-drinking rather than regular drinking). One unit of alcohol is a small glass of 13 per cent wine or a half-pint of 3 per cent lager. Consumption exceeding 14 units per week for a woman, or 21 units per week for a man, will inevitably prove harmful to health in the long term. A high percentage of the UK population, including many teenagers, currently exceeds these recommended limits.

The distinction between alcohol-related problems and alcohol dependence (addiction) is very useful. Alcohol dependence syndrome consists of:

- Withdrawal symptoms – tremor, sweating, retching and anxiety
- Relief drinking – drinking alcohol specifically to avoid or reduce withdrawal symptoms, perhaps in the morning
- Tolerance – requiring ever-increasing quantities of alcohol to achieve the same effect
- A stereotyped pattern of drinking taking precedence over other activities
- Craving
- Rapid reinstatement after abstinence, i.e. immediately resuming heavy drinking after a period of abstinence

Alcohol-related problems should be systematically sought and catalogued in the past medical history, past psychiatric history and social history.

Medical problems include gastrointestinal irritation and bleeding, cirrhosis, epileptic fits, head injuries, accidents, fractures, osteoporosis, gynaecomastia, testicular atrophy, neuropathy, pancreatitis and diabetes mellitus.

Psychiatric problems comprise anxiety, panic attacks, agoraphobia, dysphoria, deliberate self-harm, delirium tremens, amnesic black-outs, alcoholic hallucinosis, morbid jealousy, Wernicke’s encephalopathy, amnesic syndrome and dementia.

Social problems involve debt, dismissal from accommodation, work and relationships, drink-driving offences, shoplifting and domestic violence.

The features of alcohol dependence can develop insidiously in the absence of any alcohol-related problems in some people, classically in the wealthy professional who comfortably affords the alcohol, and is generally well nourished and well supported. Conversely, it is possible to accrue several alcohol-related problems in a single evening of heavy drinking with no alcohol dependency at all.

**ALOPECIA**

Barry Monk

Hair loss has a psychological impact out of all proportion to its physical significance, but disorders causing hair fall may also sometimes be a marker for systemic disorders. Convenient clinical division of the possible causes of alopecia can be made by considering: (i) whether or not obvious scalp skin abnormality is present (Table A.2); and (ii) the distribution of hair loss, for example localized, generalized or male-patterned.

**PATCHY HAIR-THINNING/BALDING ACCOMPANIED BY OBVIOUS SCALP SKIN DISEASE**

Hair loss is surprisingly uncommon in eczema and psoriasis of the scalp, even when they are severe. Allergic contact sensitivity to hair dye is a common

<table>
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<tr>
<th>Table A.2 Characteristics of alopecia</th>
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<tr>
<td><strong>Characteristic</strong></td>
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<tr>
<td>Patches of hair thinning/balding</td>
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<tr>
<td>Allergic contact dermatitis</td>
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<tr>
<td>Tinea capitis</td>
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<tr>
<td>Bacterial folliculitis</td>
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<tr>
<td>Lupus vulgaris</td>
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<tr>
<td>Morphea</td>
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<tr>
<td>Radiotherapy</td>
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<tr>
<td>Lupus vulgaris</td>
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<tr>
<td>Pseudo-pelade</td>
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</table>

| **Characteristic** | **Scalp skin abnormal** | **Scalp skin normal** |
| Diffuse hair thinning/balding | Telogen effluvium: 3 months after trigger event | Androgenic alopecia |
| Anagen effluvium: drugs and poisons | Endocrinopathy | |
| Male-patterned hair thinning/balding | Endocrinopathy | |

Patches of hair thinning/balding

Seborrhoeic dermatitis

Allergic contact dermatitis

Tinea capitis

Bacterial folliculitis

Lupus erythematosus

Lichen planus

Morphea

Hot-combing

Radiotherapy

Lupus vulgaris

Pseudo-pelade

Alopecia areata

Secondary syphilis

Trichotillomania

Traction alopecia

Alopecia totalis

Telogen effluvium: 3 months after trigger event

Anagen effluvium: drugs and poisons

Endocrinopathy

Androgenic alopecia
cause of a severe eczematous eruption of the scalp, face and neck, but hair loss is rarely a major feature. Small infants with a severe generalized atopic eczema may produce a patch of alopecia at the occiput through habitual rubbing of the head on the pillow.

A child presenting with one or more localized bald areas on the scalp associated with broken stubby hairs and scaling of the affected area of the scalp is always suggestive of tinea capitis (scalp ringworm) (see SCALP AND BEARD, FUNGUS AFFECTIONS OF, p. 588). The degree of surrounding inflammation and scaling is very variable, and depends on the fungus responsible, and the host response. Cattle ringworm (Trichophyton verrucosum) may produce a particularly violent reaction, with swelling, discharge and local lymphadenopathy, a condition termed a ‘kerion’; direct microscopy of plucked hairs and subsequent culture on Sabouraud’s medium will confirm the diagnosis. Bacterial folliculitis, if extensive enough, sometimes perpetuated by infestation with head lice, can cause patchy hair loss, sometimes with scarring developing in late or neglected cases. Pustules should be easily found, and there will be draining lymphadenopathy. A sterile inflammatory folliculitis (folliculitis decalvans) is a rare cause of patchy balding in those who are middle-aged.

**Scarring alopecia**

Various inflammatory conditions of the scalp can result in destruction of hair follicles by a scarring process; in such cases, hair loss is inevitably irreversible in the affected areas. Examination of the surface of the scalp with a hand lens reveals loss of follicles, and sometimes several hairs emerging together from a single orifice (Fig. A.9). Discoid lupus erythematosus and lichen planus are common causes of scarring alopecia. More esoteric causes include sarcoidosis (Fig. A.10), radiotherapy (Fig. A.11), lupus vulgaris and pseudo-pelade. If the scarring is linear, especially if it extends to the forehead and has a violaceous edge, localized scleroderma (morphoea) may be the cause. The whole lesion has the appearance of an exaggerated scar – en coup de sabre.

**PATCHY HAIR-THINNING/BALDING WITH NORMAL UNDERLYING SCALP SKIN**

Alopecia areata (Fig. A.12) is the most common cause of patchy baldness. Patches are asymptomatic and are often discovered by relatives or hairdressers. Patients of any age are affected, especially those in late childhood or early teens. The hallmark of this disease is a neat, sharply localized patch of billiard-ball baldness with no

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**Figure A.9** Lichen planus with scarring alopecia (Graham–Little syndrome).

**Figure A.10** Alopecia secondary to sarcoidosis.

**Figure A.11** Alopecia secondary to radiation.
obvious inflammation or scaling at the edge of lesions, and the diagnostic exclamation mark hairs should be searched for. There are usually two or three patches, and sometimes these coalesce at an alarming rate and may even cause *alopecia totalis* of the scalp (Fig. A.13), or *alopecia universalis* in which beard and all body hairs are lost. The course and prognosis are highly variable but generally good. On average, two or three patches appear, remain stable for anything up to 6 months, and then regrow without trace within 12 months. The regrowing hairs are initially often white. The cause is unknown, although there is a family history in 30 per cent of cases, and it is occasionally associated with autoimmune diseases such as vitiligo, thyroid disease, pernicious anaemia or Addison’s disease. A convincing preceding history of emotional shock is given by a proportion of patients, and may be a triggering factor. Syphilis, although relatively uncommon, is a diagnosis that must not be overlooked. Patchy alopecia may be a feature of the secondary phase. The appearance is of an asymptomatic patchy ‘moth-eaten’ baldness. On examination, there is no scaling or obvious scalp disease and, in contrast to alopecia areata, baldness is partial rather than complete. Exclamation mark hairs are not seen, and the patches are more numerous and accompanied by fever, sore throat and lymphadenopathy. The serology is positive, and the hair regrows after antibiotic treatment. *Trichotillomania* is the rather cumbersome title given to what often amounts to only a ‘habit tic’. If hair is twirled between the fingers, it eventually breaks, leaving patches of shortened hairs. Microscopic examination reveals obvious fractured ends to the affected hairs. Some psychiatrically disturbed individuals pursue hair-pulling and produce bald patches. The fractures may be seen at the scalp surface, or even at the roots. *Traction alopecia* is seen at the hair margins and is due to regular hairdressing techniques, for example rollers, braiding, ethnic plaiting and tight pony tails, pulling on the hairs (Fig. A.14).

**DIFFUSE ALOPECIA WITHOUT SCALP DISEASE**

**Telogen effluvium**

A growing (anagen) hair has a large bulb, easily seen with a hand lens on plucking. When growth ceases, the bulb shrinks, and the hair enters a resting (telogen) phase for 3 months before falling (catagen). In healthy adults, some 50–100 hairs enter telogen daily, and
thus fall some 3 months later. Not surprisingly, certain events upset the hair cycle, whereupon a larger number of hairs cease growing and enter telogen. Three months later, they will fall as a so-called ‘telogen effluvium’.

Trigerring events include childbirth, stopping the contraceptive pill, a febrile illness, blood loss, an operation, myocardial infarction, stroke, rapid weight loss, bereavement or other psychological stress. The patient often complains of a worrying increase in hair fall, but on examining the scalp, no obvious abnormality is seen although, if the hair is gently grasped between thumb and finger, many telogen hairs may be detached. Further evidence can be obtained by asking patients to collect their daily hair fall from hair brushes and pillows. Normally, some 50–100 hairs can be collected, and 300–400 can fall daily in telogen effluvium. The prognosis is excellent.

Anagen effluvium
Fall of growing hairs also causes diffuse hair-shedding, and may occur after exposure to certain drugs or poisons, for example cytotoxics, isotretinoin, thiouracil, anticoagulants, excess vitamin A and thallium poisoning.

Diffuse hair fall occurs in endocrinopathy, for example myxoedema, hypopituitarism and hypoparathyroidism. Myxoedema is regularly accompanied by hair-thinning. The mechanism is unknown and may not be directly related to serum thyroxine level, as adequate replacement therapy may fail to reverse the process. Hair loss may be a feature of systemic lupus, and it may even be the presenting symptom.

Male-pattern baldness without obvious scalp disease
Male-pattern baldness is not a disease, but an accelerated physiological process, especially pronounced in those with a genetic predisposition. Males and females progressively lose androgen-dependent scalp hairs with increasing age – in males with successive thinning of the bitemporal, occipital and pate areas, and in females with a more diffuse patterned thinning over most of the vertex. Some individuals have increased sensitivity of their hair follicles to normal levels of circulating androgens, and lose their androgen-dependent hair earlier. Such hair fall does not occur in those who have been castrated, and oestrogens and anti-androgenic drugs appear to have a protective effect. The prognosis for regrowth is poor, although many individuals search in vain for a cure.

| Table A.3 Memory nomenclature

<table>
<thead>
<tr>
<th>Suggested name</th>
<th>Immediate memory</th>
<th>Recent memory</th>
<th>Remote memory</th>
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<tbody>
<tr>
<td>Alternatives</td>
<td>Primary, short-term, working memory</td>
<td>Secondary memory</td>
<td>Semantic memory (not absolutely synonymous but conceptually similar)</td>
</tr>
<tr>
<td>Method of bedside assessment</td>
<td>Digit span test up to seven digits (repeating after no delay)</td>
<td>Recall for three to five items after 5 minutes</td>
<td>Personal history, vocabulary, beliefs</td>
</tr>
</tbody>
</table>
beliefs, and is conceptually rather similar to long-term memory. Procedural memory is outside conscious awareness, and it allows the patient to remember how to perform tasks, for example driving or cycling. It may be relatively resistant to disease processes that profoundly affect the recent memory system, such as Korsakoff’s syndrome or Alzheimer’s disease.

FUNCTIONAL ANATOMY OF MEMORY

Functional imaging of cerebral blood flow suggests that the prefrontal cortex is important for tasks involving working memory. Recent memory function involves a pathway that includes the hippocampus and the adjacent entorhinal cortex, which are richly connected to multimodal neocortical association areas. The hippocampus is thought to form new associations between ordinarily unrelated events, and damage therefore impairs learning. Midline structures, such as the medial and anterior thalamic nuclei and mamillary bodies, are also critical for recent memory. Functional imaging studies show that the hippocampus is activated during encoding; furthermore, material that evokes the most parahippocampal gyral activation is most likely to be remembered. There are anatomical links between the hippocampal formation and the midline structures, but the interaction between these structures is not well understood. The bilateral representation of the midline structures critical for memory means that bilateral cerebral damage is usually necessary to produce a severe amnesic syndrome.

Functional links between the working memory system (involving the prefrontal cortex) and recent memory system (involving the hippocampus, parahippocampal gyri and midline structures) must be important in creating long-term memories, which are likely to be stored in the neocortex.

The cholinergic neurotransmitter system plays a key role in recent memory, as shown by the damage to forebrain cholinergic projections in Alzheimer’s disease. Furthermore, cholinergic antagonist drugs, for example scopolamine, markedly impair recent memory and learning.

The synaptic basis for the encoding and storage of memories is an area of active research. The process of long-term potentiation (the modification of a synapse’s strength by the neural traffic across it) has been the most widely cited mechanism by which neural networks ‘learn’.

Memory disorders are common. When making a diagnosis in a clinical setting, it is useful to divide them according to whether the onset is rapid or gradually progressive, and whether they are of short duration or persistent. These types of memory loss are now considered.

RECENT MEMORY LOSS OF RAPID ONSET AND SHORT DURATION

Transient global amnesia

This is the prototype syndrome of recent memory loss with preserved attention. It occurs in middle-aged and elderly patients who develop sudden amnesia and bewilderment lasting several hours. There is amnesia for the recent past, as well as anterograde amnesia. They typically ask questions about their circumstances over and over again: ‘Where am I?’, ‘How did I get here?’, ‘What time is it?’ There is no impairment of consciousness, and the ability to do even complex tasks (procedural memory) is preserved. Patients remain capable of high-level intellectual performance throughout. Normal memory function will return within minutes to hours, and the patient has no subsequent recall for the period of amnesia and a brief spell before the attack. Most patients suffer only a single attack, but there is an annual risk of recurrence of about 5 per cent. The cause of this syndrome is uncertain, but antecedent events are commonly identified, including emotion or stress, cold water exposure, sexual intercourse and mild head trauma. It has been suggested that transient global amnesia (TGA) is due to an unusual form of complex partial seizure activity or cerebral ischaemia. Recent data from diffusion-weighted magnetic resonance imaging have shown restricted diffusion in the left mesial temporal lobe in seven out of ten patients during an attack, suggesting that TGA may have similarities with the cortical spreading depression thought to underlie migrainous aura propagation. A history of migraine is often found in patients with TGA.

In clinical practice, the important conditions to be considered in the differential diagnosis of TGA are complex partial seizures (which are shorter and involve altered awareness and other characteristic features – see FITS AND CONVULSIONS, p. 205), and posterior circulation ischaemia (which will usually cause additional brainstem symptoms and signs). Transient ischaemic attacks (TIAs) involving isolated ischemia of the thalamus or hippocampi may produce selectively impaired recent memory and a TGA-like syndrome. Once the diagnosis of TGA is secure, the patient can be reassured that the condition is notably benign, with no increased risk of ischaemic stroke.

Ictal amnesia

Amnesia for the duration of the seizure is usual in tonic–clonic seizures, complex partial and absence
seizures, due to disrupted electrical activity in components of the brain memory systems. There may be brief retrograde amnesia prior to attacks as well as a period of post-ictal amnesia. Memory loss may occasionally be the only symptom of an epileptic seizure involving temporal lobe structures, although observers usually describe speech or motor disturbance, or automatic behaviours. The brief episodes of memory disturbance seen in childhood ‘petit mal’ absence may cause problems with learning and behaviour. Rarely, complex partial seizures in adults may result in prolonged non-convulsive status epilepticus, which may last for days or weeks and for which the patient is subsequently amnesic.

**Electroconvulsive therapy**
Temporary impairment of memory is almost invariably following electroconvulsive therapy (ECT). It may be retrograde as well as anterograde. Unilateral ECT has much less effect on memory than bilateral ECT.

**PERSISTENT RECENT MEMORY LOSS**
The disorders in which recent memory is persistently impaired are listed in Box A.2 and will now be briefly outlined.

**Korsakoff’s syndrome**
Korsakoff’s syndrome, first described between 1887 and 1891, is a dramatic example of the amnesic syndrome. It is related to thiamine deficiency and commonly associated with long-term alcohol abuse, although it can also result from other causes of thiamine deficiency such as persistent vomiting (including hyperemesis gravidarum), intestinal obstruction, malabsorption, puerperal sepsis and metastatic carcinoma. It usually follows or accompanies Wernicke’s encephalopathy, which is characterized by confusion, ophthalmoplegia and ataxia. The definition of a pure Korsakoff’s syndrome requires that the patient is awake and attentive, responsive, and capable of understanding language, making appropriate deductions and solving problems. Newly presented information is correctly registered, but cannot be retained for more than a few minutes (anterograde amnesia or learning failure). There may be an associated variable dysfunction of recall of older memories – days, weeks or even years – i.e. retrograde amnesia. Confabulation, or falsification of memory, is commonly (but not invariably) seen. If recovery occurs, the period of retrograde amnesia shrinks but leaves a gap in memory for the period of anterograde amnesia following the onset of the illness. Neuropathological studies have shown a degeneration of neurones and loss of myelin in the mamillary bodies, the anteroventral and pulvinar nuclei of the thalamus, and the fornix.

**Head injury**
A severe head injury, sufficient to impair consciousness, invariably results in amnesia for the period of unconsciousness. It is also apt to cause retrograde amnesia, which extends for seconds, minutes or sometimes hours prior to the injury, and post-traumatic amnesia (PTA), which extends for days, weeks or, rarely, months after the injury. PTA is associated with reduced orientation and difficulty learning, and therefore has a major impact on rehabilitation. The duration of the retrograde amnesia will tend to shrink with time, whereas the anterograde amnesia is more persistent. The duration of PTA is of considerable value in assessing the severity of injury and prognosis: the longer the PTA, the more severe the head injury and the poorer the prognosis. As a guide, of patients with PTA of less than an hour, 95 per cent can be expected to return to work within 2 months; if the amnesia lasts over 24 hours, only 80 per cent will return to work at 6 months. The most severely injured may remain permanently disabled. Patients who have recovered consciousness may appear capable of conversing and carrying out normal activities, yet are unable to recall these activities later when recovery is complete because they are still in a state of PTA. This can impair their rehabilitation, and must be taken into account. Following recovery from PTA, patients may be forgetful and may complain of problems with memory for 2 or 3 years. A residual defect remaining this long is likely to be permanent.

<table>
<thead>
<tr>
<th>Box A.2</th>
<th>Causes of persistent recent memory loss</th>
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<tr>
<td>• Korsakoff’s syndrome</td>
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<tr>
<td>• Head injury</td>
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<td>• Hypoxia post cardiac arrest</td>
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<tr>
<td>• Anterior cerebral artery aneurysm rupture</td>
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<tr>
<td>• Cerebral infarction</td>
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<td>– Hippocampi</td>
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<td>– Medial thalamic nuclei</td>
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<tr>
<td>• Herpes simplex encephalitis</td>
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<td>• Limbic encephalitis</td>
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<tr>
<td>• Structural lesions of hypothalamic–mamillary body region</td>
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<tr>
<td>– Tumours</td>
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<tr>
<td>– Granulomatous disease, e.g. sarcoidosis</td>
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<tr>
<td>• Dementias</td>
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<tr>
<td>– Vascular dementia</td>
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<tr>
<td>– Alzheimer’s disease</td>
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<tr>
<td>– Dementia with Lewy bodies</td>
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<tr>
<td>• Frontotemporal dementias</td>
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<tr>
<td>• Huntington’s disease</td>
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Assessment of memory loss after head injury is difficult, and is sometimes influenced by litigation. Formal psychometric assessment of memory function should always be undertaken, although this may be difficult or impossible in the context of profound PTA. Head injuries that do not cause loss of consciousness are unlikely to result in severe amnesia. Penetrating wounds of the head, unless they specifically injure the medial temporal lobes, are also unlikely to cause problems with memory. Permanent memory defects may follow single severe acute head injuries or repeated minor traumas, as in the case of boxers (dementia pugilistica). The pathology of memory loss after closed head injury varies. Trauma can result in cerebral oedema followed by infarction of the hippocampus and cingulate gyri. Memory loss may be due to diffuse microscopic injuries causing diffuse axonal injury. Figure A.15 demonstrates burr holes to treat extensive extradural haemorrhage in a young footballer.

Vascular disease
Bilateral limbic structure infarction (including the hippocampi and medial thalamic nuclei) may cause persistent amnesia. There are often associated neurological signs to indicate posterior cerebral artery territory infarction, including visual disturbances, cortical blindness, aphasia or alexia. Unilateral infarction in the same areas may rarely cause problems with memory. Isolated frontal infarcts have also been reported to cause memory impairment. Patients who suffer rupture of an anterior communicating artery aneurysm, or undergo surgical treatment for such a lesion, may suffer ischaemia (due to vasospasm), and consequent infarction in the distribution of the small penetrating branches of the anterior communicating artery. This results in damage to the posterior inferior medial frontal areas, and to the anterior portion of the fornix and corpus callosum. These patients may present with acute amnesia, which may recover in those in whom the ischaemia is temporary and related to vasospasm.

An acute hypoxic cerebral insult, such as that resulting from cardiac or respiratory arrest, or after carbon monoxide poisoning, may produce an irreversible amnesic syndrome because of involvement of the medial temporal lobes and thalamus.

Encephalitis and other inflammatory conditions
Herpes simplex encephalitis is a striking cause of an acute persistent amnesic syndrome. Patients with this severe illness typically present with seizures, behavioural change, encephalopathy, dysphasia and hemiparesis; because of the predilection of the virus to cause haemorrhagic infarction in the temporal lobes, there may be a specific amnesic syndrome. If memory deficits persist for 1 month or more, the prognosis for recovery is likely to be poor. In addition to herpes simplex infection, any pathological process involving the functional networks underlying memory systems, particularly limbic structures, can cause amnesia. Subtle cognitive decline frequently occurs in multiple sclerosis and, in rare cases, there may be specific and severe memory impairment.

Neurosarcoidosis, cerebral lupus and neurological Behçet’s disease may also cause memory impairment. In patients with small-cell lung carcinoma, there is an associated form of ‘limbic encephalitis’ in which memory defects occur as a non-metastatic, distant manifestation of the cancer. Specific antibodies to neuronal components (most commonly anti-Hu antibodies) may be identified in serum or cerebrospinal fluid. More rarely, this syndrome can be associated with other tumours, including carcinoma of the testis or breast.

Cerebral tumour
Amnesic syndromes are rare as the presentation of cerebral tumours. They do nevertheless occur with masses arising in the diencephalus–mamillary body region in the midline. Causes include corpus callosum tumours (e.g. astrocytoma) arising in the region of the fornix. The fornix may be damaged after removal of a colloid cyst of the third ventricle, causing postoperative amnesia.

Memory loss associated with dementias
Insidious recent memory loss is the most common presenting symptom in Alzheimer’s disease, and
it becomes increasingly severe as the condition progresses. Other neurodegenerative conditions, including the frontotemporal dementias, may also involve memory function, although recent memory is typically preserved for longer into these illnesses than in Alzheimer’s disease. Dementia with Lewy bodies, progressive supranuclear palsy and corticobasal degeneration may all involve progressive recent memory impairment, but these should have other neurological features to suggest the correct diagnosis. Vascular dementia is another common cause of progressive (classically ‘stepwise’) memory impairment, and infarctions in the thalamus or hippocampi, or in the white matter pathways connecting these regions to the neocortex, are the probable cause. In all of these conditions, the progression of memory loss is usually associated with intellectual, perceptual, linguistic, praxic, attentional, personality and mood disturbances, indicating the diffuse evolving nature of the underlying pathology (Fig. A.16).

OTHER TYPES OF MEMORY LOSS

Drugs
Many drugs impair memory as part of a central nervous depressant effect, but others have a more specific amnesic effect. The latter include cannabis, organic solvents, heavy metals such as lead and mercury, anticonvulsant drugs, anticholinergic drugs and benzodiazepines. Older anticonvulsant drugs, particularly phenytoin and the barbiturates, have marked effects on memory in normal volunteers and in patients with epilepsy. The new anticonvulsant topiramate may also cause mental slowness and verbal learning disturbance. Other new anticonvulsant drugs, including gabapentin and lamotrigine, appear to have fewer cognitive side effects than older medications.

‘Psychogenic amnesia’
Complaints of memory impairment are common in depression and anxiety, but formal assessment with psychometry will usually reveal that reduced attention motivation or low mood is the cause for the symptom. More florid psychogenic amnesic states do occur, but differ from organic amnesia in the pattern of the memory defect and in the time course of onset and recovery. Loss of personal identity is common in psychogenic amnesia, but extremely rare in organic amnesia. The common setting of the ‘psychogenic fugue’, in which the patient is discovered wandering, often a long distance from home, is associated with loss of personal identity and amnesia. There may be

Figure A.16 (a) Axial magnetic resonance image (MRI) of a patient with dementia secondary to neurosyphilis, showing a generalized reduction in brain volume. (b) Coronal MRI of the patient in (a).
a triggering event such as financial or marital problems. Recovery of normal learning and alertness is often sudden, but loss of personal identity and profound retrograde amnesia may persist, unlike the usual temporal memory gradient and gradual recovery seen in organic amnesias. Inability to recognize their spouse or partner is also typical. The retrospective forgetting of circumscribed periods from the past is often found after distressing events, as in wartime, but may include periods of alleged criminal activity in malingerers. Feigned amnesia may be detected by the ‘two-choice’ recognition test of memory, in which malingerers will score significantly worse than they would by chance.

ANGIOMAS AND TELANGIECTASIA

Barry Monk

An angioma is a proliferation of blood vessels and occurs as a developmental or an acquired vascular abnormality. Telangiectasia (Fig. A.17) is the term applied to skin lesions composed of a network of fine visible blood vessels in the skin; it may arise in a number of congenital and acquired disorders.

DEVELOPMENTAL VASCULAR ABNORMALITIES

Vascular birthmarks

Transient, small salmon-pink macular birthmarks – naevi flammei – are remarkably common, and are thought to occur in over 50 per cent of live births, affecting the sexes equally. They are most commonly found on the nape of the neck, forehead and eyelids. Those on the face usually resolve within months, but a naevus flammeus on the nape of the neck more often persists into adult life.

The most important distinction that must be made in children is between a vascular naevus and a haemangioma. Vascular naevi, most commonly arising from a developmental anomaly of the dermal capillaries, are known as port wine stains (Fig. A.18). They are present from birth, and persist throughout life, growing in proportion as the child grows, and tending to darken in adult life. Not infrequently, they have a dermatomal distribution and, when arising in relation to the trigeminal nerve, may be associated with ipsilateral vascular anomalies of the brain (Sturge–Weber syndrome), which may manifest itself with fits, mental retardation or spasticity. Arteriovenous malformation (Fig. A.19) presents with pulsatile lesions, which may bleed torrentially following injury. By contrast, the strawberry naevus is a haemangioma that is absent at birth and appears in the early weeks of life. Its alarming rate of growth may be disconcerting to the parents, but spontaneous regression will follow by the age of 8 years, and active intervention is only required if the lesion interferes with the visual axis or with feeding. Rapidly growing strawberry naevi may ulcerate, and this may be associated with haemorrhage (Fig. A.20). Rarely, a massive cavernous haemangioma may sequestrate platelets and lead to a bleeding tendency (Kasabach–Merritt syndrome). Multiple haemangiomas are especially common in very premature babies. Recently the beta-blocker propranolol has been found to be highly effective in promoting the spontaneous resolution of symptomatic lesions.
Hereditary haemorrhagic telangiectasia (Osler–Weber–Rendu syndrome) is a common genetic condition manifested by multiple small vascular lesions in the skin, associated with mucosal lesions. Cases commonly present with recurrent epistaxis, or with bleeding from the gastrointestinal tract, and female patients may suffer from menorrhagia. Occasionally, there are associated vascular anomalies in the lungs; occasionally pulmonary hypertension may arise. Generalized essential telangiectasia may be distinguished by sparing of the mucosae, but the body is more widely affected with telangiectases, which are arborizing rather than spider. Ataxia–telangiectasia (Louis–Bar syndrome) is a recessively inherited immunodeficiency syndrome. Affected children are small of stature, and develop progressive cerebellar ataxia from the age of 2 years; telangiectases appear on the bulbar conjunctivae, ears and cheeks from the age of 3 years.

Acquired vascular abnormalities

Cherry angioma (Campbell de Morgan spots) develop on the trunks of almost all individuals past middle-age. They are usually small, from 1–3 mm in diameter, bright red, globular and soft. They are of no systemic significance, but are said to involute spontaneously should the eighth decade of life be reached. Larger cavernous lesions, especially on the lower lips, are common in old age (venous lakes). Small angiomas surmounted by a variable amount of hyperkeratosis (angiokeratoma) (Fig. A.21) are common on the scrotum (angiokeratomas of Fordyce), but also occur scattered in the bathing trunk area in the extremely rare Anderson–Fabry disease (alpha-galactosidase deficiency) (Fig. A.22). This X-linked recessive disorder is a condition in which the diagnosis is often delayed due to the inconspicuous nature of the angiokeratomas, but it is important to recognize because renal and vascular involvement can lead to early death.
Pyogenic granuloma has a characteristic morphology, growing on a stalk surrounded by a collarette of normal skin. These rapidly growing angiomas are seen on the chest and extremities of young people and, because of their tendency to bleed, are often the cause of alarm. A *glomerus tumour* (glomangioma) also occurs on the extremities, often beneath a nail, and is composed of a bluish-red, rounded firm papule a few millimetres in diameter. Lesions can be excruciatingly painful on pressure.

*Kaposi’s sarcoma* is a form of angiosarcoma that, in its classical form, grows indolently on the extremities of elderly Jewish or Southern Italian persons. An endemic form, more aggressive and metastasizing, was described in younger people in subequatorial East and Central Africa in the 1950s. The *epidemic* of aggressive Kaposi’s sarcoma seen in the last 30 years is largely associated with HIV infection.

*Acquired telangiectasia* is common. Isolated spider naevi (Fig. A.23) appear on children’s faces, and during late pregnancy over half of mothers develop several scattered over the face, upper chest, arms and hands. These usually disappear within 6 weeks of delivery. Similar lesions appear in *thyrotoxicosis* and *liver disease*, and also in two conditions where vasodilatory agents are intermittently released into the circulation – *carcinoid syndrome* and *systemic mastocytosis*. Other cutaneous manifestations of chronic liver disease include palmar erythema, leuconychia and clubbing. Telangiectasia on exposed skin is related to the gradual disappearance of support tissue that occurs with age, and more particularly with cumulative sun exposure. Similar mechanisms cause telangiectasia after *X*-radiation (Fig. A.24) and following the excessive use of *topical corticosteroids*. They are also seen in localized skin disorders such as *rosacea* and *poikiloderma*, as well as in collagen-vascular disorders such as *scleroderma* (matt telangiectases), *dermatomyositis* and *lupus erythematosus*.

**ANORECTAL PAIN**

*Harold Ellis*

Where there is an evident cause, the history of anorectal pain is usually of relatively short duration, and treatment is frequently successful in relieving symptoms. A small subgroup exists, however, in which symptoms are longstanding and no organic cause is found; these patients present a major therapeutic challenge to the clinician.

**CLASSIFICATION OF MAJOR CAUSES**

**Acute causes**
- Anal fissure
- Perianal haematoma
- Herpes simplex infection
- Perianal abscess
- Intersphincteric abscess

**Chronic causes**
- Proctalgia fugax
- Coccydynia
- Idiopathic
- Sometimes associated with descending perineum syndrome
- Gynaecological disorders
- Anorectal malignancy
- Presacral tumours or cysts
- Cauda equina lesions
- Tumours

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**Figure A.23** Spider naevus.

**Figure A.24** Telangiectasia after *X*-irradiation.
• Trauma
• Anal fistula
• Chronic perianal sepsis
• Crohn's disease
• Anorectal tuberculosis

SHORT HISTORY OF PAIN
Acute disorders in the perianal region usually give rise to severe pain because of the profusion of sensory nerve endings prevalent in the squamous epithelium at and below the level of the dentate line. A sudden onset of pain in association with a dark blue oedematous perianal swelling are the characteristic features of a perianal haematoma, which is thrombosis of a large venous dilatation in the external venous plexus (Fig. A.25). A history of anal pain initiated by defecation and lasting for a variable period up to an hour afterwards is usually diagnostic of an acute anal fissure. The lesion is observed on inspection of the anus usually in either the anterior or posterior midline position, and may be associated with an oedematous 'sentinel' skin tag at its more caudal margin (Fig. A.26). Digital examination or instrumentation of the anal canal causes severe pain and tenderness associated with marked spasm of the internal anal sphincter; and should be avoided at all cost. Chronicity or multiplicity of a fissure observed in unusual sites around the circumference of the anal canal should arouse suspicions of underlying Crohn's disease.

Infection with herpes simplex virus is extremely common and may present with pain due to anal lesions. Lesions are typically shallow ulcers that crust over and heal within days to weeks; tender enlargement of the inguinal lymph nodes during an attack is typical. The frequency of recurrent attacks is very variable; they affect the same anatomical site. The diagnosis of herpes simplex should be confirmed by a swab for viral culture.

The association of a short history of pain with fever and purulent anal discharge usually signifies perianal sepsis. The primary source is usually an infected anal gland and, if the sepsis remains localized, an intersphincteric abscess is the result. The diagnosis can be notoriously difficult because there may be no overt signs of infection; exquisite tenderness on digital examination of the anal canal may be the only physical finding. Usually, pus in the infected anal gland extends to the surface (i.e. to the perineum or buttock), in which case a fistula opening will be clearly visible, and an area of induration corresponding to the fistula track will be palpable.

Figure A.25 (a) Perianal haematoma, a particularly large example. (b) The clot evacuated under local anaesthetic, with immediate relief of pain.

Figure A.26 Acute anal fissure. The edges of the anal verge are gently retracted by the examiner’s fingers to reveal the fissure in the 6-o’clock position. The skin tag (‘sentinel pile’) is seen at its inferior position.
PAIN OF CHRONIC DURATION

Patients with chronic perineal pain may be found to have organic disease, although, after exhaustive investigation, no cause is apparent in many of them. Proctalgia fugax is a common source of perineal pain in which no structural abnormality is apparent. The pain is spasmodic, with episodes lasting up to 30 minutes, and is probably the consequence of paroxysmal contraction of the levator ani musculature. Coccydynia is a rather loose term applied to a history of vague tenderness and ache in the region of the sacrum and coccyx. Sometimes the pain radiates to the back of the thighs or buttocks and is usually provoked by sitting. Symptoms, without any convincing evidence, have been considered to arise from the coccyx. Idiopathic perineal pain is sometimes associated with the descending perineum syndrome, a disorder of the pelvic floor in which the pelvic floor becomes denervated, and on examination the perineum is seen to ‘balloon’ well below the bony pelvis as represented by the level of the ischial tuberosities. The pain in these patients may arise from stretching of the pudendal nerves, or alternatively from the mucosal prolapse that occurs secondarily to loss of muscle tone. Characteristically, the pain is provoked by prolonged standing or walking and is relieved by lying flat.

Of the treatable underlying disorders, malignancy in the rectum or anus must be excluded early on by digital examination and sigmoidoscopy. Gynaecological and presacral pathology should be excluded by pelvic examination, ultrasound and computed tomography scanning of the pelvis. If the history of pain accompanies a motor disorder of the anorectum and bladder, a cauda equina lesion should be suspected and excluded by magnetic resonance imaging. Finally, chronic perianal sepsis should always suggest a possible inflammatory disorder such as Crohn’s disease or anorectal tuberculosis.

ANXIETY

Andrew Hodgkiss

Anxiety is a universally experienced emotion, the presentations of which require an understanding of both its adaptive role and the relationship between personality and coping.

Anxiety is the emotional component of the ‘fight or flight’ reaction – the physiological response to threat. In ‘fight or flight’, the individual has both a choice to make and to prepare for taking either option. Once fight or flight is enacted, anxiety subsides, its job done; thus, anxiety is the emotion of indecision or conflict and of preparation for action. Anxiety contrasts with fear, which is the emotion of a non-conflictual, known threat and it precedes depression, the emotion of loss, which develops when the consequences of the threat become evident. In daily living, anxiety enhances both physical performance (e.g. in sport) and mental performance (e.g. in examinations). Being anxious occasionally makes the difference between surviving or not, but often facilitates action to ward off potential threats such as illness, pain, helplessness, punishment and separation, as well as threats to one’s status or social functioning.

How the anxiety that is generated to deal with a potential menace – whether internal or external, real or imaginary – is disposed of depends upon the individual’s personality attributes, and especially the array and efficiency of mental defence mechanisms that have been developed. To simplify, the ‘normal’, mature, balanced personality deals with anxiety effectively, either through initiating appropriate actions or through the utilization of various defence mechanisms that produce an adaptive response.

An example would be a student who feels anxious about a romance and settles their uncertainty or sublimes this emotion by intensifying their study. The immature or disordered personality cannot handle anxiety – either because the correct, relieving actions are not undertaken, or because a dearth, imbalance or inadequacy of defence mechanisms leads to a maladaptive response. In such a case, the student who feels anxious about a romance may take an overdose or dissociate from this emotion by becoming severely depersonalized and so be unable to work.

Defence mechanisms are therefore unconsciously operated mental tricks for disposing of anxiety that might otherwise overwhelm the individual. At least 30 mechanisms have been described, and both heredity and upbringing are involved in determining their presence and application. Common examples are regression, repression, denial, rationalization, projection and introjection. It follows that many types of psychological problem can develop from the failure to manage anxiety – from the interaction of personality and stress, and from the combination of coping strategies and defence mechanisms that are employed.

It should be evident that excessive amounts of anxiety may be generated in spite of a sound personality if the stress is great enough. This is the origin of post-traumatic stress disorder – how the ordinary individual copes with the extraordinary event. Excessive stress is also the basis of the adjustment reaction, a common phenomenon in medical practice when patients face the threats and uncertainty of illness – cancer or not, transplant or not, survival...
or not. These reactions tend to follow the pattern of the illness, being florid and severe in acute, life-threatening illness, and persistent and less severe in chronic disabling disorders. In adjustment reactions, anxiety figures predominantly in the early stages, and depression latterly, as the uncertainty becomes certainty and the loss apparent. Sometimes there is an unwelcome tendency for any overt emotional response to be interpreted as pathological and so to be treated with drugs: in an adjustment reaction, the best approach is enquiry and explanation, exploring the patient’s concerns, reassuring when appropriate, expediting investigations and treatment when appropriate, being a good listener. Indeed, this is an excellent opportunity to forge a therapeutic alliance with the patient that facilitates not only the recognition and prevention of further emotional distress, but also the patient’s coping with developments, setbacks and even mistakes as the illness unfolds.

It is when anxiety is inappropriate in degree or duration that the basis for diagnosing generalized/chronic anxiety disorder is established. The diagnosis of generalized anxiety disorder rests upon: (i) excessive anxiety and worry; (ii) other clinical features (Box A.3); and (iii) the exclusion of underlying physical or mental disorders.

The features of anxiety are direct consequences of increased activity of the autonomic nervous system. The symptom pattern varies considerably from patient to patient, but physical symptoms are frequently presented as the primary complaint, with anxiety and other psychological concomitants interpreted as a secondary response.

Any feature in Box A.3 may be the presenting complaint, but particularly common are headaches, dizziness, chest pain, palpitations, gastrointestinal symptoms, tremor, fatigue and emotional upset. Of course, likely physical illnesses must be excluded, even in the overtly anxious patient, and the very act of treating the symptom seriously can be therapeutic, as anxious people usually respond to reassurance. The differential diagnosis of anxiety is lengthy and complicated by the fact that anxiety frequently overlaps other disorders. This is particularly true when considering psychiatric differential diagnoses, with panic and hyperventilation both being cause and effect. The most important practical distinction is from depression, although once again anxiety and depression frequently co-exist – particularly in the less severe forms of emotional disorder typically met in general practice. The crux is to identify major depressive states in which agitation is marked. Usually, patients with agitated depression have developed typical biological and cognitive depressive changes; however, if there is doubt, it is preferable to err on the side of misdiagnosing an anxiety state as an agitated depression than vice versa.

Medical disorders that may cause or present with physical or psychological manifestations of anxiety are numerous (Box A.4). Within this list, there are several conditions that merit further discussion. Hyperthyroidism is justifiably the best-known differential diagnosis. Symptoms that point towards hyperthyroidism are increased cold tolerance, increased appetite and significant weight loss, while distinguishing signs (in addition to the classical findings in the eye and neck) are warm extremities and fine (versus coarse) finger tremor. Excluding thyroid disorder is also desirable when no obvious stressor can be established and there is no evident predisposition to anxiety.

Drugs – both stimulants (especially caffeine) and abstinence from alcohol or benzodiazepines – are another common factor: a detailed inquiry into drug-taking habits and recent changes is essential, and this should include coffee and tobacco consumption.

In general, an anxiety state arising without adequate explanation in a middle-aged or elderly person is highly suspicious of either an underlying physical disorder or a depressive illness.

Finally – and central to the whole issue of diagnosing anxiety disorders – it is usually not the illness that is masked but the doctor who is blind. By confining enquiries to physical systems, the correct questions are not asked, and consequently the correct diagnosis is missed. Perhaps the doctor may justifiably feel sometimes that a psychological line of inquiry could upset the patient and might damage their relationship, but there are usually ways round this – either by

<table>
<thead>
<tr>
<th>Box A.3</th>
<th>Clinical features of anxiety</th>
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</thead>
<tbody>
<tr>
<td><strong>Somatic</strong></td>
<td><strong>Psychological</strong></td>
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<tr>
<td>Breathlessness</td>
<td>Tension headache</td>
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<tr>
<td>Palpitations</td>
<td>Musculoskeletal aches</td>
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<tr>
<td>Accelerated heart rate</td>
<td>Restlessness</td>
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<tr>
<td>Sweating but cold and clammy</td>
<td>Coarse tremor</td>
</tr>
<tr>
<td>Dry mouth</td>
<td>Trembling, shaking</td>
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<tr>
<td>Lump in throat</td>
<td>Psychological</td>
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<tr>
<td>Nausea</td>
<td>Hyperalertness</td>
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<tr>
<td>Butterflies in the stomach</td>
<td>Feeling keyed-up, on edge</td>
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<tr>
<td>Diarrhoea</td>
<td>Feelings of dread or threat</td>
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<tr>
<td>Urinary frequency, hesitancy, urgency</td>
<td>Irritability</td>
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<tr>
<td>Dizziness</td>
<td>Increased sensitivity to stimuli</td>
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<tr>
<td>Faintness</td>
<td>Poorly sustained concentration</td>
</tr>
<tr>
<td>Light-headedness</td>
<td>Inability to relax</td>
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<td></td>
<td>Initial insomnia</td>
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</tbody>
</table>
exercising tact and judgement as to how far and fast to make the approach, or by speaking to and involving relatives. Undue delay or inappropriate referrals, investigations and treatments aggravate rather than ameliorate anxiety states, and hence make the task of helping the patient more difficult.

### Box A.4 Causes of anxiety

<table>
<thead>
<tr>
<th>Most common</th>
<th>Less common</th>
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<tbody>
<tr>
<td>- Normal/stress related</td>
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<tr>
<td>- Adjustment reaction</td>
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<tr>
<td>- Generalized anxiety disorder</td>
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<tr>
<td>- Drugs</td>
<td></td>
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<td>- Stimulants: cocaine, amphetamines, caffeine</td>
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<td>- Sympathomimetics</td>
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<td>- Hallucinogens</td>
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<td>- Drug withdrawal</td>
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<tr>
<td>- Alcohol</td>
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<tr>
<td>- Physical disorders</td>
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<tr>
<td>- Hyperthyroidism</td>
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<td>- Hypopylaemia</td>
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<td>- Hyperventilation</td>
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<td>- Migraine</td>
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<td>- Temporal lobe epilepsy</td>
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<td>- Brain tumour</td>
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<tr>
<td>- Head injury</td>
<td></td>
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<tr>
<td>- Psychiatric disorders</td>
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<tr>
<td>- Depression</td>
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<td>- Mania</td>
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<td>- Schizophrenia</td>
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<td>- Phobias</td>
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<tr>
<td>- Hysteria</td>
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<td>- Malingering</td>
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<tr>
<td>- Obsessive-compulsive disorder</td>
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<td>- Post-traumatic stress disorder</td>
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<td>- Panic</td>
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<td>- Depersonalization</td>
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<tr>
<td>- Hypochondriasis</td>
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<td>- Neurological disorders</td>
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<tr>
<td>- Cerebrovascular disease</td>
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<td>- Encephalitis</td>
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<td>- Neurosyphilis</td>
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<tr>
<td>- Multiple sclerosis</td>
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<td>- Wilson's disease</td>
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<td>- Huntington's disease</td>
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<td>- Gastrointestinal disorders</td>
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<td>- Hiatus hernia</td>
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<td>- Peptic ulcer disease</td>
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<td>- Ulcerative colitis</td>
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<td>- Crohn's disease</td>
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<td>- Cardiovascular disorders</td>
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<td>- Congestive cardiac failure</td>
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<td>- Chronic respiratory failure</td>
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<td>- Cardiac arrhythmias</td>
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<tr>
<td>- Myocardial infarction</td>
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<td>- Hypertension</td>
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<tr>
<td>- Anaemia</td>
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<tr>
<td>- Emphysema</td>
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<td>- Asthma</td>
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<td>- Autoimmune disorders</td>
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<tr>
<td>- Systemic lupus erythematosus (SLE)</td>
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<td>- Polyarteritis nodosa</td>
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<td>- Temporal arteritis</td>
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<tr>
<td>- Toxicological disorders</td>
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<td>- Benzene and derivatives</td>
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<tr>
<td>- Carbon disulphide</td>
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<tr>
<td>- Mercury</td>
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<td>- Arsenic</td>
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<td>- Lead</td>
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<td>- Organophosphates</td>
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<td>- Other causes</td>
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<td>- Vitamin B12 deficiency</td>
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<tr>
<td>- Pellagra</td>
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<td>- Aspirin intolerance</td>
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<td>- Brucellosis</td>
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<tr>
<td>- Infectious mononucleosis</td>
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<tr>
<td>- Carcinoid syndrome</td>
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<td>- Systemic malignancy</td>
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<td>- Porphyria</td>
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<td>- Uraemia</td>
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<td>- Electrolyte disturbances</td>
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<tr>
<td>- Anaphylaxis</td>
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<td>- Premenstrual tension syndrome</td>
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### LOSS OF APPETITE

Loss of appetite (or anorexia) is so common and non-specific that its presence is rarely of assistance in making a diagnosis. It can be a feature of many physical or psychological disorders, as well as a transient phenomenon in stress or even ordinary living. When a patient complains of diminished appetite, a useful pointer to the importance and clinical significance is the presence and amount of accompanying weight loss. Without confirmed weight loss or other evidence of illness, it is inappropriate to pursue investigations of loss of appetite.

Gastrointestinal disorders that are characteristically associated with loss of appetite include the prodromal stage of viral hepatitis, gastric carcinoma, gastric ulcer and coeliac disease. In coeliac disease, however, the patient may occasionally compensate for the malabsorption with an increase in appetite, and in these circumstances, loss of weight is not a problem. Patients with roundworm infestation may also have a loss of appetite but, uncommonly, may have an increase in appetite.

Anorexia may be a prominent feature of any chronic disease such as advanced malignant disease, chronic alcoholism, uraemia, severe congestive heart failure, chronic pulmonary disease and cirrhosis of the liver. In advanced cancer, profound weight loss (cachexia) can be disproportionate to a reduced calorie intake. In contrast to starvation, this cachexia cannot be reversed by simply providing more calories.

Adrenal insufficiency is often associated with anorexia and loss of weight and few other symptoms. On the other hand, both thyrotoxicosis and diabetes mellitus may lead to a marked loss of body weight in the absence of any impairment of appetite.

Anorexia may feature prominently in patients with psychiatric illness, including anxiety and depression. Anorexia nervosa is a specific psychological condition characterized by a very low body weight (BMI <18.5 kg/m² and often <17.5 kg/m²) as a consequence of a restricted calorie intake due to an intense fear of gaining weight and a distorted perception of body weight. The term ‘anorexia’ is actually a misnomer as patients often retain their appetite and have an Obsessional preoccupation with food. The condition can affect 5–10 per cent of adolescent girls and young women in some societies, with a significant morbidity and mortality. It is rare for these syndromes
to occur in males. Anorexia nervosa usually begins in a teenage girl who is either overweight or believes herself to be so. There are many accompanying physical abnormalities in the patient with established anorexia nervosa. These include amenorrhoea, osteoporosis, abnormal temperature regulation, bradycardia and hypotension, decreased glomerular filtration rate, renal calculi, oedema, constipation, and abnormality of liver biochemistry. The patient will become anaemic with leucopenia and thrombocytopenia.

**INCREASED APPETITE**

An increase in appetite will occur normally in individuals exercising strenuously, and transiently in those recovering from an illness. An increased appetite can occur in mania and hyperthyroidism. Hypoglycaemia – such as occurs with an insulinoma – may be associated with an increased appetite, but this is an uncommon manifestation of the disease. Patients with clinical depression or mania may eat to excess.

**Bulimia nervosa** is characterized by recurrent episodes of binge eating followed by inappropriate behavior to prevent weight gain – typically self-induced vomiting or laxative abuse. Appetite is usually increased, and unlike in anorexia nervosa, weight is normal or increased. The patient is usually aware that there is an abnormal eating pattern, but fears that she will not be able to stop voluntarily. After the eating binge, the patient becomes depressed. Physical features in bulimia include menstrual abnormalities, hypokalaemia, acute gastric dilatation, parotid gland enlargement, dental enamel erosion, the risk of Mallory–Weiss tears and aspiration pneumonia.

Some medicines increase appetite as a side effect. Megesterol acetate (a progestogen), dexamethasone and some antidepressants (e.g. mirtazepine) have all been used specifically to increase appetite in patients with cachexia from different causes. Drugs that cause appetite suppression as a side effect include metformin, bupropion and topiramate.

**PERVERTED APPETITE (PICA)**

Pica can occur at any age but is most common in childhood. Affected individuals repeatedly eat non-food substances such as earth or clay (geophagia), starch (amylophagia), ice (pagophagia), hair or paper. Pica may also occur during the course of pregnancy and is of no special significance. It may be associated with iron-deficiency anaemia or may form part of a mental disorder, such as autism, intellectual disability or schizophrenia.

**APRAXIA**

David Werring & Mark Kinirons

Apraxia is impairment in the execution of learned (or skilled) movements not caused by weakness, paralysis, incoordination, extrapyramidal disorders or sensory loss. One model of the execution of skilled movements is that the (dominant) left temporoparietal cortex is where visual, auditory and somatosensory information is integrated to form the motor programs for the skilled movement of both hands. These engrams then pass to the left premotor cortex, before being transferred to the right hemisphere premotor cortex via the anterior corpus callosum. Thus, lesions in the dominant temporoparietal cortex, the anterior corpus callosum or either premotor cortex can cause apraxia, and deficits of praxis can be predicted from the above model. Apraxia can be divided into three main types (affecting progressively ‘higher’ elements of the motor system) – kinetic (limb), ideomotor and ideational – as well as a number of other specific disorders of praxis.

**CLASSIFICATION**

**Limb kinetic apraxia**

This is a loss or degradation of the components of motor programs, resulting in coarse or unrefined movements that no longer have the expected appearance of being practised over time. These apraxias are usually contralateral to a hemisphere lesion.

**Ideomotor (ideokinetic) apraxia**

This is the most common type of apraxia. There is a loss of the volitional ability to perform novel or meaningless movements. A lesion in the dominant hemisphere, in the premotor area, anterior corpus callosum or inferior parietal lobe, or dementias involving the posterior hemispheres (e.g. Alzheimer’s disease) can cause ideomotor apraxia.

**Ideational apraxia**

Ideational apraxia is an impairment of ideational (conceptual) knowledge resulting in a loss of the links between tools and their actions, as well as the ability to sequence movements correctly. Movements may be left out, produced in the wrong order, or attempted with incorrect tools. It may result from diffuse brain disease or dominant parietal lobe lesions.

**Apraxia of speech**

Some patients who have disturbances of motor speech are considered to show an apraxia of buccolingual movements, although this may co-exist with, and be...
difficult to distinguish from, aphasia. The region of
damage involves the insula in the dominant frontal
lobe.

Dressing and constructional apraxia
In patients with lesions of the non-dominant parietal
lobe, there may be an inability to put on clothes in the
correct order, or to identify items that have been turned
inside out. Such patients will also show constructional
apraxia (i.e. they will have difficulty in copying a
diagram, drawing a clock or constructing a figure from
matchsticks). This disorder is probably best considered
as a type of visuospatial dysfunction rather than a true
apraxia.

Orofacial apraxia
Orofacial apraxia may be seen with lesions that
affect the dominant supramarginal gyrus or motor
association cortex. Patients will be unable to lick their
lips or pretend to blow out a match, although they will
be able to perform these tasks if there is jam on their
lips or a lighted match is held in front of them.

Conduction apraxia
Conduction apraxia causes selective difficulty in
copying movements, so that patients can pantomime
a sequence on command better than they can imitate
it. This is similar in concept to conduction aphasia (see
SPEECH, ABNORMALITIES OF, p. 625). The location of
the lesion is not known.

CLINICAL ASSESSMENT OF PRAXIS
Apraxia should be considered when the patient has
functional disability that seems out of keeping with the
degree of motor impairment shown on neurological
testing. Ideational apraxia may be suggested when
components of a complex task can be performed
individually but not integrated together as a whole.
Some simple suggestions for bedside testing are
summarized in Box A.5.

Apraxias are unusual in that the problems are often not
evident to the patient and are easily overlooked by the
examining doctor. The special types of testing required
to fully appreciate apraxias may be extremely difficult
because of co-existing neurology, such as aphasia or
agnosia.

DIFFERENTIAL DIAGNOSIS
Comprehension disorders, movement disorders
and language disorders (aphasias) must be carefully
evaluated before a diagnosis of apraxia is made.
Impaired object recognition (visual agnosia) must be
ruled out if patients select the wrong tool for a task.
Patients must be able to name the tool or describe its
function and purpose in order to establish that they
have apraxia and not visual agnosia.
The main causes of apraxia are listed in Box A.6.
It will be clear that apraxia is a set of clinical
syndromes; it is not a specific diagnosis in itself. The
history of onset and associated physical signs should
point towards the correct underlying diagnosis. For
example, stroke is usually of sudden onset without
progression, in contrast to the subacute onset of
deficits due to a mass lesion, and the slower
progression of symptoms due to a degenerative
condition. In clinical practice, it is unusual to see
apraxia in isolation. Thus, in corticobasal ganglionic
degeneration, progressive supranuclear palsy,
fronto-temporal dementias and Alzheimer’s disease,
there will be associated neurological features; these
may include bradykinesia and rigidity, supranuclear
gaze palsy, altered behaviour and memory loss,
respectively.

| **Box A.5 Bedside testing of praxis** |
|-----------------|------------------------------------------------|
| **Limbs**       | Ask the patient to: |
|                 | • Mime use of common tools, e.g. comb, toothbrush, pen |
|                 | • Imitate use of comb, toothbrush, pen |
|                 | • Use comb, toothbrush, pen |
| **Orofacial**   | Ask the patient to: |
|                 | • Whistle |
|                 | • Stick out tongue |
|                 | • Blow cheeks out |
|                 | • Cough |
| **Serial actions** | Ask the patient to mimic a serial task, e.g. ‘Mime each step, from the beginning, of how you clean your teeth.’ The patient should then be observed to mime the following: |
|                 | • Open tube of toothpaste |
|                 | • Put toothpaste on brush |
|                 | • Brush teeth |
|                 | • Rinse mouth with water |
|                 | • Spit out water and rinse toothbrush |

| **Box A.6 Differential diagnosis of apraxia** |
|-----------------|------------------------------------------------|
| **Stroke**      | |
| **Traumatic brain injury** | |
| **Cerebral tumour** | |
| **Neurodegenerative conditions** | |
| – Corticobasal-ganglionic degeneration | |
| – Progressive supranuclear palsy | |
| – Alzheimer’s disease | |
| – Frontotemporal dementias | |
| – Multiple system atrophy | |
and which may be accompanied by changes in skin paraesthesiae is usually attributed to nerve root or lesion of the brachial plexus and peripheral nerves is included, as well as cervical spondylosis.  

Lesions of the neck  
- Disc prolapse  
- Spondylosis  
- Syringomyelia  
- Fracture dislocations  
- Post-herpetic neuralgia  
- Radiculitis – paralytic/viral (neuralgic amyotrophy)  
- Spinal abscess  
  - Tuberculous  
  - Brucella  
  - Pyogenic  
- Epidural abscess  
- Pachymeningitis cervicalis  
- Tumours  
  - Spinal cord  
  - Meninges  
  - Nerve roots  
  - Vertebrae  
  - Primary  
  - Secondary  

Lesions of the brachial plexus  
- Cervical rib  
- Malignant infiltration  
- Costoclavicular compression  
- Subclavian aneurysm  
- Scalenus anterior syndrome

Lesions of the thorax and thoracic spine  
- Cardiac ischaemia  
- Syphilitic aortitis  
- Thoracic disc  
- Oesophagitis  

Lesions at the shoulder  
- Periarthritis/capsulitis  
- Subacromial bursitis  
- Calcific tendinitis  
- Bicipital tendinitis  
- Shoulder-hand syndrome  

Lesions at the elbow  
- Epicondylitis  
- Olecranon bursitis  

Lesions of the forearm, wrist and hand  
- Carpal tunnel syndrome  
- Tenosynovitis  
- Ulnar neuritis  
- Trigger finger  
- Algodystrophy  
- Hypertrophic osteoarthropathy  
- Pachydermoperiostitis  
- Repetitive strain injury, e.g. writer’s cramp

**ARCUS CORNEALIS (ARCUS SENILIS)**

**Mark Kinirons**

Arcus cornealis is an extremely common and asymptomatic bilateral peripheral corneal condition associated with hypercholesterolaemia and/or ageing. It is a type of degeneration that is found in 60 per cent of people between the age of 40 and 60 years. It is more common in black than other races and increases in frequency with age.

The opacity (Fig. A.27) is due to a deposition of lipid droplets in the superficial and deep layers of the cornea, forming a yellowish-white ring about 2 mm in width, with a clear space between it and the junction of the cornea and the sclera (the limbus). It may also arise from hyaline degeneration of the lamellae and cells of the cornea. Some individuals with corneal arcus due to hypercholesterolaemia may merit lipid-lowering treatment according to their absolute levels of cardiovascular risk.

**ARM, PAIN IN**

**Toby Garrood**

This section deals primarily with pain referred into the arm from the neck and thorax. In addition, pain arising in the brachial plexus and peripheral nerves is included, as also are lesions at the shoulder, elbow, wrist and hand that specifically or characteristically affect the upper limb. The causes of such pain are summarized in Box A.7.

Lesions that may arise at any site, such as arthritis, bone tumours, injuries and skin disease, are excluded. Pain referred to the arm falls into two major categories. Sharp, well-localized neuralgia often associated with paraesthesiae is usually attributed to nerve root or trunk compression. Dull diffuse discomfort in the limb, which is often difficult for the patient to describe, and which may be accompanied by changes in skin temperature, vascularity and sweating, is often ascribed to the involvement of autonomic pathways. In the case of this ‘cylindrical’ limb pain, an origin within the thorax or the thoracic spine should be considered.

**LESIONS IN THE NECK**

X-ray changes of cervical spondylosis are a normal finding after the age of 40. Over the age of 60, neurological symptoms and signs referred from the cervical roots are common. Great care must be taken, therefore, before ascribing patients’ symptoms to spondylosis.  

*Cervical spondylosis* can produce three clinical syndromes, which may occur alone or in combination. The first is pain and stiffness of the neck, which is often recurrent, and may be aggravated by tension, anxiety and posture. The second syndrome is radicular pain radiating down one or both arms, which may or may not be associated with muscle wasting, weakness and reflex changes referred to as cervical neuralgia. Third, there may be compression of the cervical cord, which may produce three sets of symptoms and signs:

- Weakness, wasting and fasciculation in the upper limbs, with a reduction or loss of the tendon reflexes at the level of the compression

**Figure A.27** Opacity seen in arcus cornealis.
Paraesthesiae in the arms and legs, with or without impaired sensation in the hands and feet

Pyramidal involvement with weakness, spasticity, hyperreflexia and extensor plantar responses in the legs

The combination of weakness and wasting in the arms and spastic weakness in the legs resembles amyotrophic lateral sclerosis; spondylosis may usually be distinguished from this by the history of paraesthesiae, evidence of sensory impairment, and radiographic or magnetic resonance imaging (MRI) evidence of cord compression. L'Hermite's sign may be demonstrable.

Disc herniation at the C5/C6 and C6/C7 intervertebral spaces is a common cause of pain in the upper limb. The onset may be acute, with well-localized pain radiating from the back of the neck, across the back of the shoulder, and down the arm and forearm to the wrist or fingers; more commonly, the onset is less dramatic, often after a period of recurrent aching and stiffness in the neck. Pain may be aggravated by movements of the neck, by downward pressure on the head, and by changing position of the arm. Pain may radiate downwards into the scapular region and to the upper chest. Sensory disturbances are common, although they may be detected in a dermalomat distribution (Fig. A.28), and muscle weakness may be detected in the appropriate muscles. The clinical signs associated with the most common root lesions are indicated in Table A.4. Depression of the biceps jerk may indicate a lesion of the C5 root; paraesthesiae in the thumb and index finger with depression of the supinator jerk, a lesion at the C6 root; and paraesthesiae in the index and middle fingers with loss of the triceps jerk, a lesion of the C7 root. Paraesthesiae in the feet with spasticity in the legs and extensor plantar responses indicate pyramidal damage associated with cord compression. X-rays of the cervical spine may show disc space narrowing, especially at the C5/C6 or C6/C7 levels, with lipping of the adjacent margins of the vertebral bodies. In the acute stage, X-rays may not reveal a relevant abnormality. Protrusion of a disc may be demonstrated by MRI.

Other causes of brachial neuralgia are uncommon. Viral, bacterial and fungal infections should be considered. Herpes zoster may give rise to persistent pain in the arm, especially in the elderly. The history of a vesicular rash and residual pigmented scars in a dermatomal distribution is usually diagnostic; weakness of one or more muscles in the limb with cutaneous hyperalgesia or hypoesthesia may also be present in a minority of cases.

Vertebral and paravertebral abscesses may result from tuberculosis or brucellosis, or be caused by more common pyogenic organisms such as Staphylococcus aureus. In drug addicts and immunocompromised individuals, including those with AIDS, fungal or parasitic lesions may occasionally develop. Such lesions may or may not be accompanied by fever, and the initial symptoms may closely resemble those of cervical disc prolapse. Occasionally, there are no other pointers to a septic lesion, so that severe root symptoms in the arms in the absence of clear radiographic abnormalities should prompt computed tomography (CT) or MRI examination of the neck. Pachymeningitis cervicalis hypertrophica is a rare condition, sometimes syphilitic in origin that causes diffuse pain in both arms, together with paraesthesias, widespread atrophy, loss of reflexes and variable sensory loss; more than one root is implicated.

Figure A.28 Dermatomal distribution of pain referred to the arms. (White, cervical nerve segments; grey, thoracic nerve segments.) [Redrawn with permission from Doherty, MacFarlane and Maddison (1985), Rheumatological Medicine, Churchill Livingstone, Edinburgh.]
Positive syphilitic serology should not be taken to indicate this rare condition in the absence of other diagnostic features. Primary or secondary neoplasms of the vertebral bodies may give rise to root pain with or without motor, sensory and reflex changes. X-ray examination may be diagnostic with destructive damage, although MRI is most helpful. Spurious hot spots may be seen in the presence of marked degenerative disease of the spine; plasmacytomas and myelomas produce a normal bone scan. CT and MRI scanning may be helpful in early lesions. Tumours of the meninges and roots usually cause symptoms in the legs, from compression of the pyramidal and sensory tracts, as well as pain in the arm. Root lesions in the presence of multiple cutaneous neurofibromata (von Recklinghausen’s disease) should raise the possibility of the development of a neurofibrosarcoma. Specialized spinal imaging is necessary for diagnosis. Where a neural tumour is suspected, MRI scanning will provide the most sensitive diagnostic information. Syringomyelia occasionally causes pain in the arm, but only as a late feature. By this stage, the classical features of dissociated sensory loss, muscle wasting and hyporeflexia in the arms with pyramidal signs below the level of the lesion are likely to be apparent. Fracture dislocations of the cervical spine are especially likely in the presence of rheumatoid arthritis or ankylosing spondylitis. In the former, atlanto-axial and/or subaxial subluxation of the spine may lead to upper and lower limb symptoms, and fused segments of spondylitic spine are particularly at risk of fracture with or without displacement. Fractures of cervical vertebrae due to osteoporosis are very unusual.

**LESIONS OF THE BRACHIAL PLEXUS**

Compression of the neurovascular bundle, including the brachial plexus, may occur at several sites, giving rise to characteristic features classified as thoracic outlet syndromes. Symptoms include paraesthesiae of the fingertips, especially in the night or early morning; the ulnar border of the hand is typically affected (in contrast to carpal tunnel syndrome, which affects the radial border), but numbness on waking may extend to the distal forearm. Symptoms may be aggravated by carrying heavy weights, although this is not diagnostic. Typically, pain is felt behind the clavicle and down the inner aspect of the arm, and there may be atrophy of the hypothenar eminence and intersossei. Paraesthesiae and hypoesthesia in the C8 and T1 dermatomes, with associated vasospastic features, are common findings. The diagnosis is usually based on induction of paraesthesiae and numbness by abduction of the arm to 90° with external rotation, detection of an arterial bruit in the supraclavicular fossa during this manoeuvre, and disappearance of symptoms and bruit with a return of the arm to the neutral position. Finding a position of the arm in which the radial pulse is obliterated has been considered to be a key diagnostic finding. However, this may be demonstrated in healthy subjects, and symptoms may be due to compression of the brachial plexus without involvement of the subclavian artery. The diagnosis is not, therefore, dependent on the demonstration of arterial compression. When chronic or recurrent subclavian artery compression is present, this may rarely lead to the development of aneurysmal dilatation of the subclavian artery. Causes include the position of the scalenus anterior muscle, the presence of a cervical rib, and stretching of the plexus over a normal first rib by drooping of the shoulder, which may occur in middle life. In a few patients, the accessory rib may be palpable and visible on X-ray; not infrequently, the rib is vestigial, occurring as a fibrous band that cannot be detected. In the majority of instances in which the diagnosis of thoracic outlet syndrome is considered, an alternative cause such as cervical spondylosis, cervical disc lesion or peripheral nerve lesion will be detected.

Pain in the arm is occasionally due to pressure on, or infiltration of, the brachial plexus by malignant tumours. Lymphadenopathy associated with lymphomas or carcinoma will usually be detectable by palpation of the axilla and of the posterior triangle.

---

**Table A.4 Signs and symptoms associated with common nerve root lesions affecting the arms**

<table>
<thead>
<tr>
<th>Root</th>
<th>Paraesthesia/numbness</th>
<th>Muscle weakness</th>
<th>Reflex change</th>
</tr>
</thead>
<tbody>
<tr>
<td>C5</td>
<td>Radial aspect of forearm</td>
<td>Shoulder abduction</td>
<td>Biceps jerk diminished</td>
</tr>
<tr>
<td>C6</td>
<td>Thumb and index finger</td>
<td>Wrist extension and pronation</td>
<td>Supinator jerk diminished</td>
</tr>
<tr>
<td>C7</td>
<td>Middle finger, back of hand</td>
<td>Elbow extension and finger extension</td>
<td>Triceps jerk diminished</td>
</tr>
<tr>
<td>C8</td>
<td>Little finger, ulnar border of hand</td>
<td>Finger and wrist flexion</td>
<td></td>
</tr>
<tr>
<td>T1</td>
<td>Ulnar border of forearm (see Fig. A.28)</td>
<td>Intrinsic muscles of hand</td>
<td></td>
</tr>
</tbody>
</table>
of the neck, although infiltration of the plexus by metastatic carcinoma, especially from the breast, may take a long time to become detectable. Involvement of the plexus by upward spread of an apical bronchial carcinoma (a Pancoast tumour), or more rarely by apical inflammatory lung disease, may produce unilateral Horner’s syndrome in addition to arm pain. Such lesions can usually be detected on a chest X-ray. In each of these conditions, pain may be very severe, without any accompanying signs in the early stages. Further infiltration usually leads to paralysis with relative sparing of sensation.

LESIONS OF THE THORAX AND THORACIC SPINE

In contrast to the characteristically searing localized pain of nerve root involvement, pain in the arm originating in the chest has a dull, poorly localized quality, sometimes described as cylindrical. Such pain may also be associated with alterations in autonomic functions, including temperature of the limb and sweating.

Pain associated with acute coronary syndrome (myocardial infarction or angina pectoris) is usually readily recognized, and confirmed by electrocardiography (ECG) or exercise testing. Syphilitic aortitis may induce similar referred pain. Oesophagitis may also produce cylindrical arm pain with or without more classical ‘heartburn’. Such pain may also be accompanied by ECG abnormalities, so that accurate distinction from myocardial ischaemia may rest upon cardiac enzyme analysis (troponin T and creatinine kinase), exercise testing, and visualization of the upper gastrointestinal tract. Referral from the thoracic spine is a major but little recognized cause of the upper gastrointestinal tract. Referral from the thoracic spine tenderness, often with rib and sternal tenderness, and pain on thoracic rotation. Other causes of stiffness of the thoracic spine, including spondylisis, may lead to similar symptoms. Infections of the discs also produce radicular pain. Discitis due mainly to blood-borne infection is increasingly recognized as MRI has become the investigation of choice in back imaging.

In a minority of instances, myocardial infarction leads to the development of pain and stiffness at one shoulder, with varying degrees of pain, swelling, osteoporosis and vasomotor disturbance more distally in the limb. This ‘shoulder–hand syndrome’ is discussed further in the miscellaneous conditions section.

LESIONS AROUND THE SHOULDER, ELBOW AND WRIST

In the absence of swelling, many painful lesions in the arm are referred from the spine, even in the presence of local tenderness. Thus, even apparently discrete lesions of the shoulder and elbow may originate from spinal lesions.

Degenerative arthritis at the shoulder joint is unusual. Pain around the shoulder radiating to the outer aspect of the upper arm, with pain and a reduction in glenohumeral movement in all planes, is referred to as capsulitis or periarthritis. A painful arc on abduction of the shoulder, especially with tenderness at the shoulder tip, is typical of supraspinatus tendonitis or subacromial bursitis. Transient calcification around the supraspinatus tendon may be seen on X-ray. Similarly, tenderness of the long head of biceps (bicipital tendonitis) may be noted, usually in association with capsulitis of the shoulder. The pain of tendonitis at the shoulder is usually exacerbated by resisted movement of the appropriate muscles: (i) supraspinatus – abduction; (ii) infraspinatus – external rotation; (iii) subscapularis – internal rotation; (iv) biceps – supination and flexion of the elbow.

Swelling of the olecranon bursa, due to trauma, gout, infection or other inflammatory arthritis, may produce pain over the extensor aspect of the elbow, with limitation of movement.

Nerve palsies can all cause arm pain. Inflammatory or traumatic lesions at the medial aspect of the elbow may lead to ulnar neuritis, with characteristic pain, tingling and numbness radiating down the ulnar border of the forearm and hand, with impaired intrinsic muscle function. This is especially common after prolonged bed rest, in which prolonged pressure is applied to the elbows. Radial nerve injury, producing wrist-drop with pain, numbness or tingling over the back of the hand, is more likely to result from pressure or trauma above the elbow, where the nerve runs around the posterior aspect of the humerus in the radial groove. Median nerve dysfunction produces characteristic pain, numbness and tingling in the thumb, index and middle fingers. Symptoms are often worse at night and first thing in the morning. This is most commonly caused by carpal tunnel syndrome, especially in the presence of hypothyroidism, pregnancy or inflammatory arthritis. Pressure may also be exerted on the median nerve where it passes between the two heads of pronator teres in the forearm. A positive Tinel’s sign with wasting and weakness of the abductor pollicis brevis, and slowed median nerve conduction on electromyography, confirm the diagnosis.
A variety of repetitive strain syndromes is now described. These soft-tissue syndromes relate to repeated or sustained actions of the upper limb, and produce local pain, fatigue and a decline in performance. Symptoms are most common in young adults, especially keyboard workers, and a variety of factors, including poor posture, stress, inadequate rest periods, poor training, worker’s compensation and other psychosocial factors, may contribute to their development. Both work and recreational activities may be implicated. Writer’s cramp, with pain in the wrist and shoulder associated with an excessively tight grip of the pen and a tense posture while writing, may be a related condition. Acute viral radiculitis (paralytic brachial radiculitis, neuralgic amyotrophy) produces severe pain in the shoulder and upper arm, often with a rapid onset of muscle wasting and weakness. Symptoms usually subside after a few days, although there may be some persisting weakness and ache. Tenderness over the lateral epicondyle, sometimes extending to involve the superior radio-ulnar joint, is referred to as ‘tennis elbow’, and over the medial epicondyle as ‘golfer’s elbow’. These are enthesopathies of the forearm extensor and flexor origins respectively.

MISCELLANEOUS CONDITIONS
Reflex sympathetic dystrophies may affect the upper limb, being usually referred to as shoulder–hand syndrome, causalgia or algodystrophy. This condition is characterized by pain, swelling, vasomotor disturbances and trophic skin changes usually affecting the distal part of the limb. This may follow peripheral nerve injury or myocardial infarction, although in at least 50 per cent of cases, no cause is demonstrable. In the later stages, contractures may also develop. Hypertrophic (pulmonary) osteoarthropathy (HPOA) may affect many sites but in particular the elbows, wrists and fingers. The onset of joint pain is often acute with stiffness and weakness, and there may be marked tenderness of the distal long bones associated with the radiographic appearances of periostitis. Clubbing of the fingers is also present. HPOA is usually associated with malignancy of the lung, pleura or diaphragm, although it may occasionally be benign or hereditary; it is usually bilateral and symmetrical. Only the upper limbs are affected in association with Pancoast’s syndrome and aortic aneurysms. Similar changes of periostitis and clubbing associated with thickening of the skin, especially in the scalp, may develop soon after puberty in the syndrome of pachydermoperiostosis. The condition is benign and gradually becomes inactive after a few years.

Fibromyalgia
Fibromyalgia is a syndrome of unknown cause that manifests as chronic widespread pain which includes the arms. There is an association with poor sleep patterns, cognitive symptoms, psychological co-morbidities and other somatic symptoms. Its pattern of symptoms is not consistent with a specific musculoskeletal abnormality.

ATAxia
Mark Kinirons
Ataxia is defined as uncoordinated movements that are not caused by weakness, altered tone or involuntary movements, and result from damage either to afferent sensory inputs (sensory ataxia), or to the cerebellum and its connections (cerebellar ataxia).

Cerebellar Ataxias
The cerebellum is involved in the regulation of muscle tone, the coordination of movement, and the control of posture and gait. Cerebellar dysfunction will arise with lesions of the cerebellum itself or its connections in the cerebellar peduncles, midbrain, pons or medulla (Fig. A.29). The cerebellar hemispheres relate ipsilaterally to the limbs, and therefore unilateral lesions of the cerebellar hemisphere will result in ataxia that is most marked on the same side of the body. Midline cerebellar lesions tend to manifest...
with severe gait ataxia. As well as ataxia, lesions of the cerebellum and its connections may give rise to additional signs such as hypotonia, pendular reflexes, dysarthria (due to incoordination of the muscles of voice production, leading to ‘scanning’ or ‘staccato’ speech) and nystagmus.

Limb ataxia is demonstrated using the finger–nose and heel–shin tests. An intention tremor may be present in the upper limbs, together with evidence of dysdiadochokinesis on rapid alternating movements of the upper and lower limbs. Patients with gait ataxia tend to walk with a broad-based gait, and persistently grab for support. If mild, this becomes most evident on heel–toe walking. Truncal ataxia is a feature of midline cerebellar lesions and manifests as an inability to sit or stand without assistance, and to fall backwards. Many different pathologies may result in cerebellar ataxia.

Cerebellar ataxia of abrupt onset
The sudden onset of cerebellar ataxia implies an aetiology that is vascular, demyelinating or inflammatory, most commonly due to posterior circulation stroke or multiple sclerosis (Fig. A.30).

Subacute evolution of cerebellar ataxia
The subacute evolution of an ataxic syndrome should raise the possibility of a mass lesion within the posterior fossa, most commonly a secondary. Haemangioblastomas and medulloblastomas also occur. Cerebellar abscesses are relatively rare, except in patients with evident infection in the middle ear; encephalitides are extremely uncommon. In some patients with lung cancer, there is a form of paraneoplastic cerebellar ataxia that develops subacutely and particularly affects the limbs.

Chronic cerebellar ataxia
Chronic alcoholism, severe hypothyroidism, vitamin B12 deficiency, phenytoin toxicity, anoxic injury including carbon monoxide poisoning, and head trauma can all cause chronic cerebellar ataxia.

The slow development of a cerebellar ataxia may also occur in the context of the hereditary ataxias, the best known of which is the autosomal recessive Freidrich’s ataxia. The cardinal features include gait and limb ataxia, dysarthria, areflexia, sensory loss and corticospinal tract signs. Skeletal abnormalities, diabetes and cardiomyopathy may also occur.

SENSORY ATAXIA
The term ‘sensory ataxia’ is used to describe the incoordination that is seen in patients who have lost proprioception in a limb or limbs. Afferent proprioceptive impulses travel from the muscles and joints, providing information to the cerebellum and cerebrum about the movements of muscles. In the absence of this sensory information, or if it is distorted, the corticospinal system is unable to coordinate normal movements. Such proprioceptive sensory loss may be seen in patients with peripheral neuropathies, spinal cord disease affecting the posterior columns (e.g. subacute combined degeneration of the cord due to vitamin B12 deficiency, or multiple sclerosis), and lesions of the thalamus and sensory cortex. The resulting ataxia is most apparent when visual guidance is removed. Associated signs include loss of touch and joint position sense, and pseudo-athetosis of the fingers (the patient being unable to keep the fingers still in the outstretched position).

The gait is typically wide-based, and the feet may strike the ground with extreme force, causing a stamping gait. When asked to stand with the feet together and the eyes closed, the patient will sway (Rombergism). In the upper limbs, the phenomenon may be demonstrated by asking the patient to hold the arms outstretched and to close the eyes. The patient is likely
to have great difficulty in undertaking fine movements that are outside the normal range of vision, such as fastening the top button of a shirt, or tying a tie.

**ATHETOSIS**

David Werring & Mark Kinirons

The word ‘athetosis’ derives from the Greek for ‘unfixed’ or ‘changeable’ and means slow, writhing, continuous movements often affecting the limbs, but also the axial muscles, including the neck, face and tongue. These movements are often confined to the upper limb, and are most pronounced in the fingers and hands. When absent at rest, the movements in one region can be brought out by asking the patient to voluntarily move other parts of the body; for example, speaking may induce athetosis in the limbs, neck, face or tongue. This phenomenon is called ‘overflow’.

Within the patterns of movement, it is often possible to identify extension, pronation and flexion–supination of the forearm, and flexion–extension of the fingers with the thumb often being trapped by the flexed fingers as the hand closes. In the lower limbs, there is typically eversion–inversion of the foot; in the face, there is retraction and pursing of the lips, and alternate constriction and relaxation of the forehead with opening and closing of the eyes. The neck and torso are prone to rotatory twisting movements.

In general, the movements of athetosis are slower and less jerky than those described in chorea, but gradations between the two forms of involuntary movement are seen; in some cases, the distinction is impossible, and the term ‘choreoathetosis’ is used. The most common cause of athetosis is damage to the basal ganglia in early life (neonatal or infancy) in the context of cerebral palsy.

In children, particularly in association with hemiplegia due to cerebral palsy, athetosis can be unilateral, although bilateral (‘double’) athetosis may also occur. Both the arms and face are involved; speech and swallowing are also affected. Such children are usually slow in gaining their motor milestones and have learning difficulty. Athetosis in infants usually involves slow twisting movements, and it is most appropriately considered as a form of dystonia.

Athetosis is considered a result of basal ganglia (extrapyramidal) diseases. In adults, athetosis may occur as an episodic or permanent abnormality in hepatic encephalopathy, as a manifestation of chronic intoxication with phenothiazine-based or other dopamine-modulating antipsychotic drugs, or, most commonly, as an effect of chronic administration of l-dopa in patients with Parkinson’s disease. Athetosis is part of the involuntary movement seen in patients with Huntington’s chorea, and it may also be seen in patients with Wilson’s disease, neuronal brain iron accumulation syndromes and Leigh’s disease. Adult athetosis has a speed approaching chorea, and is usually termed ‘choreoathetosis’. Athetosis is also a common feature of cerebral palsy. Rarely, it can result from focal brain lesions including stroke.

Athetosis, like most forms of involuntary movement, ceases during sleep and may be brought out at the bedside by distraction, for example by asking the patient to close their eyes and recite sequential numbers or days of the week out loud.

**AURA**

David Werring & Mark Kinirons

An aura may be defined as a premonitory symptom, which is commonly related to an epileptic attack. The term may also be used to describe the focal cerebral symptoms occurring before the development of headache (or sometimes without headache) in migraine.

One of the main differentiating features between the aura of an epileptic seizure and the aura of migraine is duration. The aura in migraine may last from a few minutes to an hour or more, although a good rule of thumb is that if the symptoms last for over 30 minutes, another cause, such as a cerebrovascular event, should be considered. In epilepsy, the aura is usually shorter, lasting from seconds to minutes.

**SEIZURE AURA**

A seizure aura has been defined as that part which occurs before consciousness is lost (or automatisms begin), and which can be remembered subsequently. The typical characteristics of an epileptic aura are that it is spontaneous, normally in the context of otherwise good health, begins abruptly, and lasts for seconds only. Auras are most commonly seen in patients who have complex partial seizures arising in a focal brain area, and reflect where the seizure starts. Thus, a motor aura may include involuntary movement of a limb, or occasionally a more coordinated movement such as running. A sensory aura is more common, and may be described as discomfort, tingling or occasionally a more complex sensation moving along a limb. A visual aura may be simple, consisting of shimmering lights or colours within the field of vision, implying an origin in the occipital lobes; or it may be a complex hallucination, such as flowers or a brightly coloured pattern, which suggests an origin in the lateral temporal neocortex.

Seizures arising in the medial temporal lobes are likely to be associated with a psychical aura, involving...
feels of apprehension, a sense of unreality or a ‘dreamy state’, consisting of vivid memory-like hallucinations, or the sense of having previously lived through exactly the same situation before (déjà vu). There are often associated epigastric phenomena (e.g. an abnormal sensation beginning in the stomach and welling up into the throat) and fear. These symptoms may progress to a loss of contact with the environment, oro-alimentary automatisms (e.g. chewing, lip-smacking, sucking or blowing movements), and then simple gestural automatisms. With involvement of the temporal lobe uncs, there may also be an olfactory or gustatory aura in which patients may describe disturbances of smell or taste, which are usually unpleasant.

The most important aspect of the aura in epilepsy is in helping to classify the seizure type, which can guide investigation and treatment. A focal aura indicates that the epilepsy is arising at a focus within the brain and is therefore a form of partial seizure. Thus, a patient who describes an aura consisting of a sensory disturbance in the left hand will be likely to have a lesion in the contralateral parietal lobe; one who describes an aura beginning in the right visual field will be likely to have a lesion in the left occipital lobe. An aura of movement beginning in the right thumb and spreading in Jacksonian fashion up the arm implies a lesion in the left motor strip (precentral gyrus of the left frontal lobe). All patients with focal aura must be investigated with cerebral imaging using computed tomography or magnetic resonance imaging if these are available. Traditionally, carbamazepine has been the first-line treatment for partial seizures.

The second importance of an aura in epilepsy is that it provides evidence of continuing seizure activity. Patients with seizures may be able to describe minor episodes in which their usual aura does not proceed to a full attack, thus indicating to the physician that, although control is partially achieved in reducing the severity of the attacks, the epileptogenic focus continues, and the patient requires an increase in anticonvulsant therapy.

MIGRAINE AURA

Migraine affects about 10 per cent of the population. Most suffer from common migraine (migraine without aura) in which headache, usually unilateral, is associated with photophobia, phonophobia and nausea or vomiting. One in ten migraineurs have classical migraine (migraine with aura), in which some or all of the attacks are preceded by symptoms that are typically positive (i.e. an excess of normal perceptions or other neurological phenomena) attributable to focal alterations in brain activity. The most common is a visual aura, classically a black and white or coloured zigzag (fortification) in part of the field of vision. Other auras include complex disturbances of function such as shivering loss of vision (‘scintillating scotomas’), aphasia, or sensorimotor dysfunction of part of the body. Zigzag visual forms are generally considered pathognomonic of migraine. The aura may evolve over a few minutes and continue (generally up to about 30 minutes, to a maximum of about 1 hour). If the symptoms are negative (i.e. a loss of function such as numbness or paralysis) or prolonged, the possibility of a vascular event including ischaemic stroke must be considered. It should be remembered that migraine can be part of a primary headache disorder but can also result (as a ‘secondary’ phenomenon) from other pathologies, particularly cerebrovascular disorders including carotid artery dissection, venous sinus thrombosis, cerebral arteriovenous malformations, and syndromes associated with anticardiolipin antibodies. Migraine with complex aura is a common feature of the rare familial disorder of cerebral autosomal dominant arteriopathy with subcortical infarcts and leucoencephalopathy (CADASIL), in which a progressive dementia and family history should suggest the diagnosis.

Migraine auras are usually – but not always – followed by a typical lateralized, throbbing headache. Aura without headache is termed acephalgic migraine, and quite commonly develops in middle to late adult life, especially in women. The diagnosis of transient ischaemic attack (TIA) may be considered with recurrent episodes of aura without headache, but TIA’s may be distinguished by an abrupt, non-evolving onset and must mimic a known vascular territory syndrome. Migrainous aura is more common than TIA at all ages. It should also be noted that the character of a migranous aura may change in an individual at different stages of their life; sometimes in women, aura changes are clearly related to hormonal changes. Because of the wide range of causes of migraine aura-like symptoms, brain imaging, including MRI with vascular sequences, is often indicated, particularly in the case of a first attack.

The most commonly accepted explanation of pathophysiology for migraine aura is spreading cortical depression that moves across cerebral areas at the same rate as symptoms evolve clinically, and this has recently been demonstrated on functional brain imaging. Similar studies indicate that the ‘neuronal generator’ for migraine is in the midline brainstem structures, including the periaqueductal...
grey matter, dorsal raphe nuclei and locus coeruleus, with subsequent activation of the trigeminal vascular system (see HEADACHE, p. 258).

**AXILLARY SWELLING**

**Harold Ellis**

**CLASSIFICATION**

The differential diagnosis of axillary swelling is best carried out by considering the individual anatomical structures present in the axilla and the lesions that can arise from these:

**Skin and subcutaneous tissue**
- Abscess
  - Acute
  - Chronic
- Accessory breast
- Lipoma

**Lymph nodes**
- Acute adenitis
- Chronic adenitis (tuberculosis)
- Lymphomas
- Secondary malignant deposits

**Chest wall**
- Cold abscess
- Tumours
  - Benign
  - Malignant

**Axillary blood vessels**
- Axillary aneurysm
  - Traumatic
  - Subacute bacterial endarteritis

A swelling in the axilla in the majority of cases is due to enlargement of the lymph nodes. An acute lymphadenitis will be secondary to infection in the cutaneous area drained by the axillary nodes, that is to say, the whole of the upper limb, the breast and the skin and subcutaneous tissues of the trunk, front and back, down to the level of the umbilicus. The whole of this area must be carefully searched in an effort to find a primary focus. Any form of tumour other than involvement of the axillary nodes by secondary deposits is distinctly rare, but unfortunately it is common to find the axillary nodes to be the site of metastases from carcinoma of the breast. Skin tumours, especially melanoma, in the cutaneous area already defined are a not uncommon source of axillary node deposits. The differential diagnosis is considered in more detail in the following sections.

**ACUTE ABSCESS**

An acute axillary abscess may be recognized by the well-marked signs of local inflammation, usually accompanied by the general features of pyrexia and malaise. The patient resists movement of the arm on account of the pain, and there is usually some cause, such as a paronychia, to account for the source of infection.

Although most examples of axillary abscess are due to *Staphylococcus aureus*, the remainder are caused by mixed anaerobes, often complicating hydadenitis suppurativa of the axillary skin. This is a chronic infection of the axillary apocrine sweat glands, with sinuses, scarring and a history of intermittent purulent discharge.

Rarely, an empyema points in the axilla (empyema necessitans). There are generally, but not always, abnormal lung signs to suggest the diagnosis, but the underlying lesion will be revealed on chest X-ray.

**TUBERCULOUS ABSCESS**

A tuberculous abscess forms a single fluctuating non-tender swelling in the axilla. It may result from breaking down of the tuberculous axillary lymph nodes, or from a cold abscess originating in one of the upper thoracic vertebrae that tracks round the intercostal bundle and results in a fluctuant swelling at any point along the chest wall, commonly in the mid-axillary line.

Occasionally, a cold abscess will result from a BCG injection administered in the deltoid region.

**AXILLARY LYMPHADENOPATHY**

If the examination proves that the swelling is not an abscess, the next most likely diagnosis is lymphadenopathy. This may present as a single enlarged node, a number of discrete individual nodes or a mass of matted glands.

For a detailed differential diagnosis, see Lymphadenopathy (p. 394). It is sufficient here to enumerate the principal causes. These are acute infection and metastatic deposits. Much less often encountered is chronic infection with tuberculosis, or one of the lymphomas.

An acute lymphadenopathy is suggested by the presence of septic focus in the drainage area of the axillary nodes, although not infrequently the primary focus (e.g. a splinter in the finger) has disappeared and can only be ascertained from the history. The story of a scratch on the hand from a cat, which again by now might have completely healed, suggests the possibility of cat scratch disease.
The most likely primary tumours to give rise to axillary deposits are carcinoma of the breast and a malignant melanoma of skin (Fig. A.31). Occasionally, a node or group of nodes in the axilla appears malignant and, upon being removed for histological examination, is found to be infiltrated with metastatic carcinoma, yet no source of the primary lesion can be found. The most likely site for such a hidden primary is undoubtedly an occult tumour in the breast, and this will be identified on mammography. Occasionally, a primary tumour will be discovered in the lung on chest X-ray.

ACCESSORY BREAST
Occasionally, a woman will present with a soft, subcutaneous axillary mass, resembling a lipoma that is, in fact, an accessory breast. The history of the swelling will go back to puberty, and careful examination will usually reveal an overlying nipple, which will clinch the diagnosis.

PRIMARY TUMOURS OF THE AXILLA
The only common tumour of the axilla is a lipoma (Fig. A.32), which may attain a large size and extend up under the pectoral muscles. The mass slowly grows over a period of years, is subcutaneous, has a definite outline, is freely mobile, is lobulated and gives the sign of fluctuation. This is not, of course, because fat is liquid at body temperature, but rather that each fat cell acts as a microcyst.

This condition may be confused with a cystic hygroma of the axilla. It usually arises in infancy or is even present at birth, and is associated with the more common cystic hygroma of the cervical region, which is brilliantly transilluminable.

Unusually, a mass on the medial wall of the axilla proves to have its origin from the underlying ribs, chondroma or bony secondary deposit.

AXILLARY ANEURYSM
An axillary aneurysm is rare, but the diagnosis should be obvious as it presents with an expansile pulsating mass along the line of the axillary artery. A bruit may be elicited on auscultation.

There is usually a preceding history of trauma, such as a stab wound, which has produced a true or a false aneurysm.

In the absence of any history of local injury, or in cases of an apparently spontaneous aneurysm, there may be the symptoms and signs of bacterial endocarditis.

Figure A.31 A huge mass of melanotic deposits in the axillary nodes following previous resection of malignant melanoma of the posterior of the upper arm. Note the skin graft at this site.

Figure A.32 A massive, but benign, lipoma of the axilla.
BACK, PAIN IN
Fred Heatley and Jonathan Lucas

(See also SPINE, TENDERNESS OF, p. 632.)

Pain in the back is one of the most common complaints in general and specialist practice. No speciality is immune from it, and the differential diagnosis therefore covers most of medicine. The classical approach with a long list of differential diagnoses separated out into the standard categories – traumatic, degenerative, inflammatory, infective, etc. – is indeed formidable, as illustrated in Box B.1. A close inspection will rapidly reveal that even this is abbreviated, and probably incomplete. Furthermore, few of us will retain this whole list in our brain each time we see a patient with backache and, if we did, we would probably never finish our day’s work!

Mechanical backache – i.e. musculoskeletal causes – accounts for the overwhelming majority of patients presenting with backache, no matter whether it be acute or chronic. The skill in dealing with this condition is to be able to recognize the small percentage of cases that present with non-mechanical backache and have a serious underlying disease. To avoid mistakes, the following ‘rules’ are helpful.

HISTORY

In taking the history, note the following:

- The patient is well and fit, has not lost weight, does not have night sweats, etc.
- The pattern of pain. Mechanical backache is nearly always episodic. Beware the patient who has severe constant pain localized to a particular site in the spine. This suggests underlying bone pathology, for example infection (tuberculosis is on a rapid increase throughout the world due to its association with AIDS) or malignancy.
- Always enquire about urinary symptoms. Central compression of the cauda equina is rare, often rather atypical in presentation and may not have associated abnormal peripheral neurology. However, if it is missed, and hence allowed to progress to bladder paralysis, it is devastating to the patient’s quality of life.
- Pay attention to the age of the patient. Men of 55 and upwards presenting with their first episode of backache that has persisted for more than 2–3 weeks should be presumed to have carcinoma of the prostate with spinal secondaries, until proven otherwise. Classical disc disease often starts in the late teens or early twenties with acute backache that only later progresses to episodes of sciatica.

The main differential diagnoses in this younger age group are: (a) spondylolisthesis, i.e. a stress fracture of the pars interarticularis; (b) an inflammatory discitis; and (c) ankylosing spondylitis. Females aged over 60 presenting with an acute onset of severe backache have a stress fracture of a vertebra until proven otherwise. Do not presume that these are all osteoporotic, as they may be metastatic. Osteomalacia is another diagnostic ‘blind spot’.

<table>
<thead>
<tr>
<th>Box B.1 Causes of backache</th>
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<tbody>
<tr>
<td><strong>Mechanical: traumatic and degenerative</strong></td>
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<tr>
<td>Low back pain; postural – fatigue – obesity; pregnancy</td>
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<tr>
<td>Fractures and fracture dislocations</td>
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<tr>
<td>Ligament sprains and facet joint injuries (although these are probably a common source of acute backache, they are unfortunately impossible to confirm, even with modern imaging techniques such as magnetic resonance imaging. This group falls into the category of acute self-limiting back pain)</td>
</tr>
<tr>
<td>Intervertebral disc lesions</td>
</tr>
<tr>
<td>Osteoarthritis of the facet joints, ankylosing hyperostosis with bony spondylolyses extending from the vertebral bodies around the discs</td>
</tr>
<tr>
<td>Spinal stenosis and nerve root impingement at the exit foramina</td>
</tr>
<tr>
<td>Lumbar instability syndromes – spondylitis, spondyloarthropathy including the degenerative type, i.e. secondary to facet joint osteoarthritis (see SPINE, DEFORMITY OF, p. 627)</td>
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<tr>
<td>Spinal deformity – scoliosis postural, structural. Scheuermann’s disease (see SPINE, DEFORMITY OF, p. 627)</td>
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<tr>
<td><strong>Metabolic</strong></td>
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<tr>
<td>Osteoporosis</td>
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<tr>
<td>Osteomalacia</td>
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<tr>
<td>Paget’s disease of bone</td>
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<tr>
<td>Hyper- and hypoparathyroidism</td>
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<tr>
<td>Ochronosis, fluorosis</td>
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<tr>
<td>Hypophosphataemic rickets</td>
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<tr>
<td><strong>Infective</strong></td>
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<tr>
<td>Ankylosing spondylitis</td>
</tr>
<tr>
<td>Spondylitis associated with Reiter’s syndrome, psoriasis, ulcerative colitis, Whipple’s and Crohn’s diseases</td>
</tr>
<tr>
<td>Polymyositis and polymyalgia rheumatica</td>
</tr>
<tr>
<td>Arachnoiditis (this low-grade inflammatory condition was a complication caused by the contrast medium used for myelography. Fortunately, this investigation has now been superseded by magnetic resonance imaging and computed tomography scanning)</td>
</tr>
<tr>
<td><strong>Surgical</strong></td>
</tr>
<tr>
<td>Osteomyelitis</td>
</tr>
<tr>
<td>Acute – pyogenic (see SPINE, TENDERNESS OF, Figs S.51 and S.52, p. 633)</td>
</tr>
<tr>
<td>Subacute – Salmonella (typhoid and paratyphoid fever)</td>
</tr>
<tr>
<td>Chronic – tuberculosis, brucellosis (undulant fever – Brucella abortus and melitensis), syphilis, yaws, Weil’s disease (Leptospirosis icterohaemorrhagica)</td>
</tr>
<tr>
<td>Intervertebral disc – pyogenic discitis,iatrogenic discitis (secondary to surgical exploration or investigations)</td>
</tr>
<tr>
<td>Spinal ‘space’ infection, acute and chronic meningitis, subarachnoid abscess, epidural abscess</td>
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</tbody>
</table>

(Continued)
On examination:

- Look for fixed deformities, especially a kyphos.
- Check the spinal movements. Mechanical disease nearly always has a full or almost full range of movement in at least one direction. For example, a patient presenting with an acute disc and a lumbar tilt/scoliosis will usually have very reasonable range of movement to the side in the direction of the tilt. A totally rigid spine with loss of movement in all directions should be presumed to have underlying serious disease until proven otherwise.

- Examine the peripheral neurology. If this is abnormal, check carefully that it fits with the diagnosis in the back. If a nerve is already ‘irritable’ due to an underlying neurological disease, it will be at risk of paralysis from much less compression than a normal nerve. For example, a patient with motor neurone disease or a peripheral neuropathy may present with a profound foot drop in association with a recurrent episode of acute backache from a ‘minor’ disc protrusion. Also beware a patient who has a definite loss of a knee jerk since this has a dual root innervation. (In theory, this is supplied by L2, L3 and L4, but it is predominantly L3 and L4.) Although a disc protrusion taking out both these nerve roots can occur, it is very uncommon. Note that in clinical practice there is no reflex for L5, and you need to test the power of big toe extension for the L5 root (extensor hallucis longus and brevis).
- Always do the Babinski (plantar) response. On the sensory side, the most important of all signs is loss of sensation in the saddle area – S2, S3, and S4, with loss of bladder and bowel control indicating cauda equina syndrome.

INVESTIGATIONS

In considering plain X-rays, it is ‘normal’ for the spine to show degenerative changes in patients aged over 60; indeed, these will be present in many patients aged over 50. It is all too easy to miss the absence of a pedicle, which is frequently the earliest X-ray change in metastatic disease. At least 50 per cent of metastatic deposits will probably not be visible on routine X-rays. In particular, the spinal X-ray may be ‘normal’ in early cases of multiple myeloma. Involvement of two bodies and the intervening disc is the characteristic feature of infection. Finally, destructive lesions in the sacrum and sacroiliac joints are difficult to spot on plain films due to presence of overlying bowel shadows. Do not neglect simple blood investigations such as haemoglobin, white count, acid phosphatase, erythrocyte sedimentation rate (ESR) and C-Reactive protein (CRP). In particular, if the ESR and CRP are normal, one is unlikely to have missed either the inflammatory or the infective group of diseases.

In summary, to diagnose mechanical disease with confidence it is necessary to have the following: a fit patient; a past history in which the backache and/or radicular pain has been episodic; a pattern of spinal movements in which a range of movement in at least one direction is largely preserved; neurological signs – if present – that fit to the mechanical signs; an X-ray that is ‘normal’ for the patient’s age group; and normal routine blood tests. Having diagnosed a mechanical
problem, beware of the patient who may have cauda equina compression. Fortunately, this is rare but, when present, symptoms can be surprisingly vague and easily overlooked by the unwary.

**MECHANICAL CAUSES**

This group accounts for the vast majority of backache. The principles of diagnosis and the pitfalls to be avoided before concluding that backache is indeed mechanical have already been set out in the preceding paragraphs.

**Disc degeneration**

The intervertebral disc is an ingenious structure. The central nucleus is highly hydrophilic and exerts high osmotic pressure. It is contained by the endplates above and below and the annulus circumferentially. It acts rather like an elastic ball, the range of movement between the endplates being constrained by the annulus, the ligaments and the shape of the facet joints. With increasing age, the nucleus desiccates, losing its ability to maintain the osmotic pressure, the annulus develops fissures, and nuclear material herniates in all directions, including superiorly and inferiorly into the adjacent vertebral bodies, which is manifest on X-ray as a Schmorl’s node. Marginal osteophytes form around the endplates and the bulging annulus to complete the picture of spondylosis seen on a routine X-ray. Posteriorly, the facet joints lose their congruity and, since they are synovial joints, they develop osteoarthritis. The nerve root exit canals become narrowed due to osteophyte intrusion and a change in shape secondary to the loss of disc space height. These pathological changes set the scene in which a nerve root entrapment in the younger patient is caused predominantly by disc protrusion, while in the older patient it is secondary to entrapment at the exit foramen. These changes are most marked in the distal lumbar segments (i.e. L4/L5 and L5/S1). The common disc protrusion is posterolateral, and will therefore catch the nerve root exiting under the pedicle of the vertebra below – i.e. the L5/S1 disc will most commonly compress the S1 nerve.

Classically, *disc prolapse* (Fig. B.1) first occurs in young adults – the 20- to 30-year age group. Initially, there is an episode of acute backache following lifting

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**Figure B.1** A prolapsed L5/S1 disc (magnetic resonance imaging scanning). (a) Sagittal scan; note the large disc protrusion into the spinal canal at the L5/S1 level. A small part of the disc has actually sequestered (has separated and is lying loose) (arrowed). (b) Axial view across the L5/S1 level. The prolapsed disc (solid arrow) is impinging not only into the spinal canal but also into the exit foramen. Note the displacement of the S1 nerve root (dotted arrow). The patient had severe right-sided sciatica with pain radiating down the back of the thigh and the back of the calf, and into the foot. The ankle jerk was diminished.
or bending. Spinal movements are restricted, and there is often a tilt. Frequently, this initial episode settles, and only with subsequent episodes does it become clear that there is nerve root compression. Pain in the lower back is poorly localized and is referred over a wide area, particularly to the region of the posterior superior iliac spine and into the buttock. Clinically, to diagnose sciatica one requires the pain to radiate down below the knee into the calf, preferably to the ankle. The straight leg raise will be limited. The presence of abnormal neurological signs is clearly helpful in localizing the nerve root under compression.

Plain X-rays are not particularly helpful in disc disease since in younger patients they are usually normal, whereas in the older patient one can expect to see signs of degeneration. Magnetic resonance imaging (MRI) is the investigation of choice. The main indication for surgical decompression is the persistence of severe leg pain, not necessarily the presence of abnormal neurology. Finally, beware of the patient who has had sciatica first down one leg and then down the other, since this may represent a central disc prolapse. Always enquire about micturition, and test for perineal ‘saddle’ anaesthesia.

The bony spinal canal is, like the cranial cavity, a ‘closed box’, and nerve compression can therefore result from any space-occupying lesion. The situation in the spinal column is complicated by the fact that the shape and size of the bony canal is different at different levels in the spine. Furthermore, the spinal cord, which ends opposite L1, is more at risk of compression than the cauda equina. It is fortunate that thoracic disc herniation is rare in clinical practice, even though small prolapses are quite frequently seen on MRI imaging, since thoracic discs have a much greater incidence of neurological complications (e.g. lower-extremity weakness, and bowel and bladder symptoms), and there may be a sensory level, while upper motor neurone signs may be present in the lower limbs.

Spinal stenosis (Fig. B.2) is seen predominantly in those aged over 55 years. (The sex distribution of the condition is more or less equal). (see also LOWER LIMB, PAIN IN, ‘Causes of intermittent and neurogenic claudication’, p. 351). Classically, it presents with neurogenic claudication. Although this is a well-recognized syndrome, the precise source of pain is poorly understood. It is easily confused with arterial claudication. Patients normally give a long history of low lumbar backache. The symptoms in the legs are varied – pain, weakness and tiredness – and these characteristically worsen on walking until they cause the patient to stop. Classically, this is aggravated by walking down a slope when the spine is extended, and eased by going up a slope when the spine is flexed. The symptoms disappear after resting. The lumbar spine is particularly affected (individuals with achondroplasia are particularly at risk due to their short pedicles). Within the normal population, there is a natural variation in the cross-sectioned shape of the bony canal, but the trefoil pattern – in which the lateral recesses are narrow – has an increased incidence. Old disc disease, facet joint osteoarthritis and bony encroachment spondylolisthesis will clearly reduce the space even further. Specific causes of narrowing include spondylolisthesis (especially the degenerative type), Paget’s disease of bone and, rarely, post-traumatic fracture dislocations. Compression at two levels has a much greater risk of precipitating neurogenic claudication than stenosis at one level. It is postulated that this is due to venous congestion and reduced blood flow in the intervening segment of the nerve root. (For further discussion, see LOWER LIMB, PAIN IN, ‘Leg pain of radicular or vascular origin’, p. 350.)

Nerve root entrapment at the exit foramen may often be aggravated by walking, and is easy to confuse with spinal stenosis. However, with entrapment, the pain tends to be constant, and is often present even at rest. Root tension signs, for example a limited straight leg raise, are usually found.

Spinal deformity

A structural scoliosis is rarely painful except when degenerative disease has superimposed later in life. Scheuerman’s disease – an increased thoracic kyphosis in adolescence – is usually painless but can sometimes produce aching discomfort. Like scoliosis, it can cause backache in later life (see SPINE, DEFORMITY OF, p. 629).

METABOLIC CAUSES

Osteoporosis is of increasing concern in many populations as the percentage of the elderly rises. There are three well-recognized orthopaedic entities – wrist fractures, hip fractures and vertebral fractures – all of which cause much morbidity and much expense for healthcare systems. The surprising feature about vertebral osteoporosis is that it is so often relatively symptom-free. The patient loses height and becomes round-shouldered as the vertebral bodies, especially the thoracic spine, collapse and wedge anteriorly. The posterior elements remain intact. For most patients, there is a slow subsidence rather than an acute collapse. When the latter occurs, there is a sudden onset of pain, often following a minor fall. This pain usually settles over a few weeks, but it
Figure B.2 Spinal stenosis, secondary to severe degenerative spondylolisthesis at the L4/L5 level. (a) Lateral X-ray showing the forward slip of L4 on L5 (arrowed). (b) Magnetic resonance imaging (MRI) sagittal cut. Note the sharp cut-off at the L4/L5 level. (c) MRI axial scan at the L4/L5 level. Note the severe diminution of the spinal canal; compare this with (d). (d) MRI axial scan across the body of L5, where the spinal canal is again patent. The dural sac is clearly outlined, as are both S1 nerve roots, which are just exiting in their Dural sheaths (arrowed). The epidural fat is clearly displayed (white).
may occasionally be associated with intercostal nerve root entrapment. This is easily overlooked and results in unnecessary suffering since this condition often responds well to injections of long-lasting local anaesthetic, which also of course confirms the diagnosis. Many patients suffer from multiple vertebral body fractures, and in severe cases the rib cage comes to rest on the iliac crest and can be a source of considerable discomfort. Fortunately, neurological problems secondary to spinal cord compression are very rare, even with multiple fractures. Patients may, however, have difficulty walking if the thoracic and lumbar deformities are so severe that they struggle to get their centre of gravity in line with their feet and consequently topple forwards.

It is important to realize that when radiologists refer to a bone being osteoporotic, they are using this purely as a descriptive term – ‘the bones appear osteoporotic’. This merely means that there is less calcium present to absorb the X-rays and therefore the radiograph appears darker. Several important and serious conditions can produce ‘radiological osteoporosis’. These include multiple myeloma, metastases and osteomalacia. Osteoporosis is best defined as ‘reduced mass of bone per unit volume of bone’. On densitometry measurements, osteoporosis is two standard deviations below the sex- and age-related mean, while osteopenia is one standard deviation below. In osteoporosis, routine blood investigations, including the ESR, are normal. The ESR is usually raised in metastatic disease and often reaches high levels of 80–100 mm/hour (Westergren) in myeloma. Marrow studies and electrophoresis will clinch the diagnosis in myelomatosis, but if there is any doubt as to the aetiology of a crush fracture, a biopsy under X-ray control should be carried out. Radiologically, beware of the crush fracture that involves both endplates, as there is a greater incidence of malignancy or osteomalacia. In osteoporotic fractures, it is usually the superior endplate that collapses down.

In osteomalacia, as in rickets, there is inadequate mineralization of bone, and unmineralized osteoid seams accumulate on the surfaces of new bone. There is often a history of dietetic and/or intestinal insufficiency, chronic disease or a previous gastrectomy – i.e. the inadequacy of calcification may be due to calcium deficiency or to a defect anywhere along the metabolic pathway of vitamin D. The serum alkaline phosphatase is often elevated, serum calcium and plasma phosphate are normal or decreased, whereas urinary 24-hour calcium output is low. Clinically, these patients present with diffuse backache, in contradistinction to the localized severe pain of a crush fracture from osteoporosis. The backache is usually eased by rest and aggravated by activity. Classically, X-rays of the spine show a biconcave appearance to the vertebral bodies – the ‘cod fish vertebrae’ – while X-rays of the pelvis may show Looser’s zones and, in some instances, protrusio acetabuli.

Hyperparathyroidism may present with generalized backache and tenderness. Radiographs show ‘osteoporosis’, the serum calcium is raised (one may need repeated estimations to confirm this), plasma phosphorus may be low (although it rises with renal failure), and the alkaline phosphatase is usually, but not invariably, raised. There may be other features of hypercalcaemia such as nausea, vomiting, muscle weakness or a true myopathy, corneal calcification (band keratitis), nephrocalcinosis and renal tract calculi. Peptic ulceration and pancreatitis may occur. The syndrome is usually due to primary hyperparathyroidism, secondary to hyperplasia or an adenoma of the parathyroid gland, but may occur secondary to renal or other diseases in which the serum calcium can be normal. The finding of plasma chloride levels consistently less than 100 mmol/l in the presence of hypercalcaemia virtually excludes the diagnosis of primary hyperparathyroidism. Finally, there may be other bone changes, including bone cysts (secondary to brown tumours of von Recklinghausen); subperiosteal reabsorption of the phalanges and distal end of the clavicles is also characteristic.

Paget’s disease, in which there is increased bone turnover involving both deposition and resorption, can be a cause of severe backache, although it is often an incidental finding. The characteristic X-ray feature is of an enlarged vertebral body with coarse trabeculation. Severe pain of a radicular origin is caused by a nerve root entrapment at the exit foramen. Paget’s disease can occur in a single bone or in many bones. An onset of pain or an exacerbation of pain in an old Paget’s bone may signify malignant change to an osteosarcoma, which carries a very poor prognosis. Skeletal pain secondary to Paget’s usually responds to calcitonin. However, the pain is often difficult to distinguish from that of osteoarthritis, which is common if the Paget’s disease involves a joint. Osteoarthritic pain will, however, not respond to calcitonin.

INFLAMMATORY CONDITIONS (SPONDYLOARTHROPATHY)

The best example is idiopathic ankylosing spondylitis. Here, the patient is usually a male aged between 16 and 36, and in 90 per cent of cases the
histocompatibility antigen HLA-B27 is found. The spine is stiffened and restricted in movement in all planes. Neck movements are often restricted (see NECK, PAIN AND/OR STIFFNESS, p. 450, Fig. N.19), and intercostal expansion at nipple level is reduced from the normal 5–7.5 cm to 2–5 cm or less. This intercostal restriction occurs early in the course of the disease; it is not a late complication but an essential and early part of the clinical picture. Diaphragmatic movement is normal. Evidence of active or old iridocyclitis is present in over 20 per cent of the patients, in most cases seen as iritic adhesions or dark spots on the posterior surface of the cornea. Tender heels or tender areas over the pelvic brim, ischial tuberosities or greater trochanters are not uncommon. Peripheral arthritis occurs in some 24 per cent of cases initially and hydrarthrosis of the knees in about 7 per cent. Occasionally, the disease presents as an inflammatory arthropathy of one of the large joints, for example the hip. It is easy to overlook fixed flexion deformity of the hip and erroneously to attribute the inability of the patient to stand upright solely to the spinal disease (Figs B.3 and B.4). The ESR is elevated in almost all cases, but sheep-cell agglutination and latex tests are negative. Nodules do not occur, and nor does lymphadenopathy or splenomegaly.

The most common initial symptom is aching in the buttocks, patients drawing their hand down the back of the buttocks and thighs at the site of discomfort, but lumbar backache and stiffness soon occur and may be the initial symptoms. Two radiographs help in early diagnosis − a posteroanterior view of the sacroiliac joint, and an anteroposterior film of the dorsolumbar spine D8–L3 − but X-ray changes may not be present until symptoms have been present for 2–3 years or more. The earliest radiological sacroiliac changes are blurring of the joint outlines with para-articular ilial sclerosis, erosions and apparent widening of the joint space, gradually giving way over the years to narrowing and obliteration of the joint. Small syndesmophytes, resembling bony ‘stalagmites and stalactites’, are usually seen, first along the edges of the intervertebral discs between the vertebral bodies of D10 and L2; this is where the ‘bamboo spine’ usually first becomes evident. Lytic lesions with periosteal elevation and ‘whiskering’ are most commonly seen in the pelvis or in the spine; girdle joints (hips and shoulders) secondarily, and peripheral joints least often, in contrast to the distribution of joint involvement seen in rheumatoid arthritis, where initial involvement is usually feet, hands and wrists. The spondylitic pattern of disease may also occasionally be seen in Whipple’s disease and Behçet’s disease and, very rarely, in polymyalgia rheumatica. Some male cases of
juvenile chronic polyarthritis progress to the spondylitic picture. In the diagnosis of ankylosing spondylitis, these variants should always be considered.

**INFECTIVE CONDITIONS**

A spinal infection is easy to overlook, and as a consequence there is often a considerable delay in diagnosis. For example, with tuberculosis it is uncommon to make the diagnosis within 6 months from the onset of backache, and a delay of a year is frequent. *Pyogenic osteomyelitis* can also be deceptive, especially when it occurs in patients who are ill, for example those in intensive care units and those who are immunocompromised, have AIDS or are drug addicts. A wide variety of organisms can be encountered, and a bacteriological diagnosis is mandatory.

The cardinal features are backache and a fever. Spinal infection is, surprisingly, a diagnostic blind spot in considering a pyrexia of unknown origin (PUO). In the early stages, the backache is often mild, albeit persistent. It is present at rest, although, since it is aggravated by activity, it is all too easy to attribute pain to a mechanical cause. In the established disease, night sweats and weight loss become prominent. Spinal infection is even harder to diagnose when there is no fever, as can happen with a low-grade infection in susceptible ill patients. The author has known cases in which a technetium bone scan, performed in desperation on account of continuing unexplained weight loss, showed a hot spot in a vertebral body. Whole-body MRI scans also occasionally discover ‘hidden infection’.

The hallmarks as regards the clinical signs are local tenderness, presence of a kyphos, muscle spasm and restricted movements in all directions. These signs are also easy to miss. Look for localized tenderness and a kyphos by examining the patient prone. The sharp angle of the kyphos is best felt by running a finger down the whole length of the spinous processes. Tenderness is best elicited by ‘springing’ of the spine (see SPINE, TENDERNESS OF; p. 632). In comparing the thoracic and lumbar spines, a kyphos is more prominent in the thoracic spine, whereas in the lumbar spine the natural lordosis merely flattens; however, muscle spasm and loss of mobility is more obvious in the lumbar spine since the thoracic spine is splinted by the ribs.

The classic radiological picture is the involvement of two bodies and the intervening disc, reflecting the embryological segmental pattern. However, there is considerable variation, and the infection may involve purely the body, the posterior elements or the disc (see SPINE, TENDERNESS OF; p. 632 Figs S.51 and S.52). The extent of bone destruction is usually much more marked than is expected on the basis of the symptoms and signs. Inflammatory markers, the ESR and C-reactive protein, are raised. A technetium bone scan can be helpful in localizing the site of the bony infection in patients presenting with a PUO. Computed tomography (CT) and MRI scans clearly delineate the extent of the bony and soft-tissue involvement. The longer the delay in making the diagnosis, the greater the risk of: (i) bony destruction causing significant deformity; (ii) the formation of abscesses; and (iii) neurological complications secondary to either compression of the cord or cauda equina dependent on the level of the lesion, or the entrapment of nerve roots secondary to compression/deformation at the exit foramina. In all cases of spinal infection, it is essential not only to examine the distal peripheral neurology but also to check perineal sensation and enquire about bladder control. A neurological deficit secondary to a compression by ‘soft tissue’ (i.e. abscess and inflammatory oedema) has a much better prognosis than when it is caused by angulation of the cord over a sharp bony prominence.

It is mandatory to establish the precise bacteriological diagnosis. This is best done by aspiration and biopsy under radiological or CT-guided control (see Fig. S.51). A bony specimen should be sent for histology as well as bacteriology. *Staphylococcus aureus* is the most common pyogenic infection, but a wide variety of other organisms can be found, for example a *Pseudomonas* infection in drug addicts. The differential diagnosis for the more chronic tubercular infection is from typhoid, both *Salmonella typhi* and *Salmonella paratyphi*, *Brucella abortus* from cattle, *B. melitensis* from goats, and *B. suis* in pigs. Weil’s disease (caused by *Leptospira icterohaemorrhagica*), which is characterized by fever, high white blood cell count, jaundice and haemorrhagic manifestations, can also cause acute backache at the time of the infection and may lead on to subsequently marked degenerative change and chronic backache.

*Primary disc space infection* occurs in children and adolescents prior to the closure of the endplates. In adults, it is nearly always iatrogenic – i.e. secondary to a procedure involving the disc (e.g. a discogram). Backache, which is often severe, is usually localized to the affected area – particularly so when a lumbar disc is involved – and there may be referred pain in the buttocks or thighs. The tenderness is localized, but the muscle spasm is usually widespread. The white cell count is often within normal limits, but the C-reactive protein and ESR are usually raised. In children, the condition was well recognized in pre-antibiotic days and was termed ‘benign osteomyelitis’, as it was self-resolving.
Initially, the X-rays are normal, but after 2–3 weeks disc space narrowing becomes apparent, and this eventually progresses to bone fusion. Primary extradural abscesses are uncommon, but when they occur they most frequently affect the thoracic spine. They cause severe pain that radiates along the relevant thoracic nerve root. Epidural infection is fortunately rare since it spreads rapidly throughout the spine, precipitating paraplegia and death.

**NEOPLASTIC CONDITIONS**

Metastatic disease is by far the most common malignant tumour found in bones. However, by comparison with the very high incidence of mechanical backache, metastatic spinal disease is an uncommon entity, and it is too easy to fail to diagnose malignancy when backache is the presenting symptom and the primary is unknown. Only a good thorough history will prevent the clinician from making this mistake, since in the early stages the physical signs are subtle – localized tenderness and muscle spasm. The cardinal features are: (i) backache that is persistent, unremitting, not relieved by rest and indeed often worse at night; (ii) weight loss; and (iii) no past history of mechanical backache. A useful rule states that a man presenting in his mid-50s with his first ever episode of **significant backache** should be presumed to have prostatic cancer until proven otherwise.

On X-ray there is considerable variation. Metastatic tumours may be sclerotic (prostate, sometimes breast) or lytic, solitary or multiple. Any part of the vertebra can be involved, and one of the classic radiological patterns is loss of a pedicle on the anteroposterior X-ray. Unfortunately, this is all too easy to overlook (see Figs S.53 and S.54). A technetium bone scan can be very helpful in defining the extent and number of metastases, and hence in defining which part of the skeleton should be closely studied by CT and MRI scanning. If the primary lesion is unknown, it is often easier for the patient, as well as for the doctor, to establish diagnosis by performing a CT-guided biopsy rather than carry out extensive medical investigations. The diagnosis can be particularly difficult when the presenting feature is a solitary metastasis, as the differential diagnosis then includes the whole range of malignant bone tumours in addition to the search for a primary. Renal carcinoma (hypernephroma) is often silent, and quite commonly presents with a destructive expansile lytic bone secondary. These are highly vascular. Indeed, if the metastasis is in a superficial bone (e.g. the skull), a defect will be not only palpable but also pulsatile.

Metastatic disease in the spine can also present with acute severe pain secondary to collapse of the vertebral body. This can be accompanied by acute-onset distal paralysis. This is a surgical emergency as the cord or cauda equina will need to be decompressed and the spine stabilized if the patient is to have any relief from the severe pain and to regain any quality of life.

**Multiple myeloma** nearly always involves the spine (see NECK, PAIN AND/OR STIFFNESS, p. 450, Fig. N.16). The skull, ribs, sternum and pelvis are other common sites. The patient is usually aged over 50 years, and men are affected twice as often as women. The ESR is usually raised, and a monoclonal protein is found on serum electrophoresis. Radiologically, there are round, punched-out lytic areas with no surrounding sclerotic margin. However, there is occasionally an altogether different picture of diffuse osteoporosis. This pattern can be easily overlooked – especially on poor-quality X-rays – and even when it is recognized, it is all too easy to dismiss it as osteoporosis. A **plasmacytoma** is a solitary form of myeloma, the majority of which (70 per cent) will progress to multiple myeloma. This solitary type carries a very good prognosis in the other 30 per cent of lesions if adequately treated.

**Malignant lymphoma** involves the skeleton in 20 per cent of cases, the bony lesions being distributed one-third to the lower limb, one-third to the spine and pelvis, and one-third to the rest of the skeleton. Primary bone lymphomas without general involvement are uncommon, but they do carry the best prognosis. Normally, they present with local pain, and as the patient is usually in good general health, the diagnosis can be difficult in the early stages. Skeletal pain is the presenting symptom in 25 per cent of children and 5 per cent of adults with **acute leukaemia**. Eventually, X-ray changes will be found in as many as 70–90 per cent of patients, but the pattern varies widely from transverse lucent metaphyseal lines in children to generalised osteopenia and osteolytic destruction.

**Spinal tumours**

Tumours of the central nervous system are an important group, particularly in children. Most are intracranial, but 15 per cent are intraspinal, of which 33 per cent are intramedullary, 22 per cent intradural and 45 per cent extradural. Symptoms are often rather vague, and localizing signs may take a long time to develop. Plain X-rays may show widening of the canal, scalloping of the vertebral bodies or enlargement of an intervertebral foramen. The most common intramedullary tumours are ependymomas and astrocytomas (Fig. B.5). In children, astrocytomas predominate and tend to involve the cervical and thoracic regions, while in adults the majority are ependymomas and involve the lumbosacral region.
Meningiomas, neurofibromas and haemangioblastomas are the most common extramedullary intradural spinal tumours seen in adults. The classic dumb-bell neurofibromas cause intra- and extradural expansion and usually arise from the spinal roots, the posterior more often than the anterior. These may be either single or multiple, and may or may not be part of the generalized neurofibromatosis. Extradural tumours, of course, include the whole range of bone tumours as well as tumours that can arise from the neural crest, of which ganglioneuromas are usually benign and neuroblastomas usually malignant. These usually present as large paraspinal masses that are visible on abdominal or chest X-rays. Among this large group of benign bone tumours and tumour-like conditions, the most common to be encountered in the spine are haemangiomas, aneurysmal bone cysts, osteoblastomas and osteoid osteomas. Even these are rare. Osteoid osteomas classically produce a painful scoliosis in a teenager or young adult. Characteristically, the pain is well relieved by analgesics (see Figs S.45–S.47). Haemangiomas can lead to extensive bone destruction, although the common X-ray pattern is for the vertebral body to show vertical sclerotic striations.

Backache can be a prominent feature in some patients during a sickle-cell crisis. Localized bleeding in the form of an epidural haematoma is a complication of spinal anaesthetics. In the early stages, the fluid content gives a high signal on the T2 image, which later changes to a low T2 signal as fluid is absorbed and haemosiderin deposited.

**REFERRED PAIN**

Backache may be caused by cardiovascular and intra-thoracic disorders, of which a good example is the intense, demoralizing, boring pain of an aneurysm invading the spine. Syphilitic aneurysm of the arch and upper descending aorta is associated with signs of an aortic reflux, collapsing arterial pulses and possibly signs of neurosyphilis. Dissecting aneurysms of the descending aorta below the arch are less apparent; unequal or delayed pulses in the arms and legs should be noted. An arteriosclerotic aneurysm of the abdominal aorta often causes pain in the lower part of the back as well as in the upper abdomen, the groin and occasionally the testicles; a pulsating mass can be felt in the abdomen. A carcinoma of the bronchus or oesophagus may cause backache.

A rare cause is an enormous enlargement of the left atrium with mitral disease. The pain in such cases is usually relieved by leaning forwards and to the left. The pain from myocardial infarction only rarely radiates to the back.

Chronic pancreatitis and carcinoma of the pancreas may cause a dull, persistent, upper lumbar ache that is usually (but not always) associated with upper abdominal pain and discomfort. The pain is eased by leaning forwards. A penetrating ulcer on the posterior wall of the stomach or first part of the duodenum quite characteristically gives a boring pain in the upper lumbar region; the pain is related to meals and may be relieved by antacid therapy. Enlargement of the liver from any cause may give a dull ache felt to the right of the lower thoracic spine, in addition to aching discomfort in the abdomen and lower chest. The pain of cholecystitis and cholelithiasis is felt posteriorly over the liver, or a little higher, in addition to the upper abdomen. Backache is commonly found in association with renal and genitourinary disease. Pyelitis and pyelonephritis can cause lower thoracic and lumbar backache. Renal tumours (in particular carcinoma; see above) often remain silent, as haematuria and the findings of a palpable mass in the flank only tend to occur late in the onset of the disease. Backache in association with prostatic metastases is all too easy to confuse with mechanical backache (see above).
Gynaecological conditions are a rare cause of low lumbar and sacral backache. It is common for all varieties of pain (including backache) to be worse during menses, but this does not equate to having a gynaecological cause. Thus, although backache may occasionally occur in association with diseases of the ovaries or Fallopian tubes, this diagnosis should only be made by excluding other causes. Backache due to retroversion of the uterus is a diagnosis of despair and should be discarded as a mistake.

PSYCHOGENIC FACTORS

Psychogenic factors are important to assess, especially in patients complaining of chronic backache. It is, however, important to remember that pain by itself is rarely a hysterical symptom, since these are much more dramatic as well as visible. The problem with pain is assessing the severity. Pain out of proportion to the underlying pathology is a common diagnostic problem. ‘Inappropriate signs’ are a helpful indicator, for example excessive spinal tenderness, a false straight leg raise, etc. Characteristically, with the former even the lightest touch to the skin of the back causes severe pain, and the patient may often wobble and lose balance. In a false-positive straight leg raise, the leg can hardly be lifted off the couch with the patient lying supine, but the patient can be deceived into sitting up to 90° with the legs fully extended. Before attributing a stacking–glove sensory loss to being psychogenic, a peripheral neuropathy must be excluded. Benign intraspinal tumours, cauda equina compression from spinal stenosis or a central prolapsed lumbar disc, a prolapsed thoracic disc, and myelopathy due to compression in the cervical spine, for example in rheumatoid disease, are all entities that can be associated with long-standing, rather vague symptoms and subtle signs. It is all too easy to label such patients as ‘hysterical’.

BEHAVIOUR, ANTISOCIAL

Andrew Hodgkiss

All societies generate long lists of behaviour that are considered antisocial. These vary over time and from place to place, and they reflect the values and prejudices of the whole community. It is a matter of great current debate how wise doctors are to have involved themselves in this domain at all. Accusations of medicalizing wrong-doing, being agents of social control or being apologists for criminals all haunt psychiatrists in particular. Yet the fact remains that a proportion of those who behave antisocially do so because of a definite psychiatric or medical disorder, and we would wish to avoid the punishment of those not responsible for their acts.

The most contentious areas are the concepts of conduct disorder in children and antisocial personality disorder in adults (discussed in the past as juvenile delinquency and psychopathy, respectively). These are syndromes defined to a large extent by the antisocial behaviour of the individual, yet are offered by psychiatrists to explain this behaviour! Conduct disorder is one of the most common diagnoses made by child psychiatrists. It refers to behaviours ranging from truancy, lying, drug misuse and defiance to fire-setting, stealing, serious assaults and deliberate self-harm. Although the testing of boundaries and challenging authority are part of the normal psychological work of adolescence, there is a growing number of young people in the UK whose antisocial behaviour is seriously damaging themselves and others (Box B.2). One fashionable theory is that the absence of adequate fathering or male role models is compensated for by extreme herd behaviour, in which groups of adolescent males form gangs and set their own norms of behaviour (which may include extreme risk-taking and offending). The child psychiatrist must exclude causes of antisocial behaviour such as emotional disorder, early-onset psychosis, learning disability and epilepsy. After that, there is a lack of consensus on how best to intervene through the criminal justice system or mental health services.

ADULT ANTISOCIAL BEHAVIOUR

Antisocial behaviour in adults tends to decrease with age. A medical or psychiatric basis should be suspected when the behaviour occurs acutely, unexpectedly or after a recent stressful life event. Apparently motiveless behaviours occurring in bursts are suspect. Perpetrators who are female, older and without a criminal record merit close attention. Assessment should include consideration of a detailed and accurate account of the behaviour itself from a reliable impartial informant, and a description of the premorbid personality and past mental health of the perpetrator. Any planning before, and actions and attitudes after, the event should be elicited. The mental state

<table>
<thead>
<tr>
<th>Box B.2 Causes of delinquency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Most common</td>
</tr>
<tr>
<td>• Normal</td>
</tr>
<tr>
<td>• Socially determined</td>
</tr>
<tr>
<td>Less common</td>
</tr>
<tr>
<td>• Neurotic</td>
</tr>
<tr>
<td>• Stress reaction</td>
</tr>
<tr>
<td>• Conduct disorder (antisocial personality)</td>
</tr>
<tr>
<td>Rare</td>
</tr>
<tr>
<td>• Organic, especially epilepsy</td>
</tr>
<tr>
<td>• Psychotic, especially schizophrenia</td>
</tr>
<tr>
<td>• Learning disability</td>
</tr>
</tbody>
</table>
antisocial behaviour beginning in adolescence

Offences related to alcohol and drug misuse have a personality of psychotic patients who offend. A rarity). Considerable attention is now being paid to the about hearing voices that obliges you to hit someone’

As a professor once remarked, ‘There is nothing of considering their responsibility for their behaviour. Famous Diagnosing a major psychiatric disorder in an

There are recognized associations between certain offences and particular psychiatric disorders: murder followed by suicide in psychotic depression; infanticide in severe postnatal depression; shoplifting and depression; morbid jealousy and wife murder; schizophrenia and matricide; psychoses (incorporating erotomanic delusions) and attacks on the famous. Diagnosing a major psychiatric disorder in an offender should be the beginning rather than the end of considering their responsibility for their behaviour. As a professor once remarked, ‘There is nothing about hearing voices that obliges you to hit someone’ (irresistible command hallucinations being, indeed, a rarity). Considerable attention is now being paid to the personality of psychotic patients who offend.

Offences related to alcohol and drug misuse have a sad familiarity and predictability, but the individual is generally held legally responsible for these. Persistent antisocial behaviour beginning in adolescence

and continuing into adulthood points to antisocial personality. To make a convincing diagnosis (if diagnosis is the right word) of antisocial personality disorder, the doctor needs to demonstrate that the patient displays a lack of remorse, a lack of empathy, a failure to learn from punishment and poor impulse control. There is usually a pattern of fractured relationships and work record, along with polysubstance misuse. There is little optimism about the treatment of this disorder outside forensic psychiatry services.

**Box B.3  Differential diagnosis of antisocial behaviour in adults**

<table>
<thead>
<tr>
<th>Most common</th>
<th>Delusional disorder</th>
</tr>
</thead>
<tbody>
<tr>
<td>No medical or psychiatric component</td>
<td>Schizoaffective disorder</td>
</tr>
<tr>
<td>Stress reaction</td>
<td>Mania</td>
</tr>
<tr>
<td>Adjustment reaction</td>
<td>Major psychotic depression</td>
</tr>
<tr>
<td>Neurological disorders</td>
<td>Antisocial personality disorder</td>
</tr>
<tr>
<td>Cerebrovascular accident</td>
<td>Tricyclic antidepressants</td>
</tr>
<tr>
<td>Encephalitis</td>
<td>Monoamine oxidase inhibitors</td>
</tr>
<tr>
<td>Neurosyphilis</td>
<td>Anticholinergics</td>
</tr>
<tr>
<td>AIDS</td>
<td>L-Tryptophan</td>
</tr>
<tr>
<td>Multiple sclerosis</td>
<td>Dopaminergics</td>
</tr>
<tr>
<td>Subdural haematoma</td>
<td>H2-receptor blockers</td>
</tr>
<tr>
<td>Head injury</td>
<td>Corticosteroids</td>
</tr>
<tr>
<td>Epilepsy</td>
<td>Narcotic analgesics</td>
</tr>
<tr>
<td>Dementia</td>
<td>Psychiatric disorders</td>
</tr>
<tr>
<td>Delirium</td>
<td>Hysteria</td>
</tr>
<tr>
<td>Alcohol</td>
<td>Malinger</td>
</tr>
<tr>
<td>Illicit drugs</td>
<td>Post-traumatic stress disorder</td>
</tr>
<tr>
<td>Stimulants</td>
<td>Other conditions</td>
</tr>
<tr>
<td>Opiates</td>
<td>Premenstrual tension syndrome</td>
</tr>
<tr>
<td>Hallucinogens</td>
<td>Diogenes syndrome</td>
</tr>
<tr>
<td>Prescribed drugs</td>
<td>Hypoglycaemia</td>
</tr>
<tr>
<td>Benzodiazepines</td>
<td>Anticholinergics</td>
</tr>
<tr>
<td>Drug withdrawal</td>
<td>L-Tryptophan</td>
</tr>
<tr>
<td>Learning disability</td>
<td>Dopaminergics</td>
</tr>
<tr>
<td>Major psychiatric disorder</td>
<td>H2-receptor blockers</td>
</tr>
<tr>
<td>Schizophrenia</td>
<td>Corticosteroids</td>
</tr>
</tbody>
</table>

### BLEEDING

Mark Kinirons

(See also BRUISES, p. 76; PURPURA, p. 554.) Within the context of a normal haemostatic system, excessive haemorrhage tends to occur secondary to a structural lesion, such as a peptic ulcer. Recurrent bleeding mainly from one site may point to the location of a pathological lesion. More generalized bleeding is seen with abnormal haemostasis, examples of which are isolated coagulation defects (e.g. haemophilia), platelet disorders (e.g. thrombocytopenia) and drug effects (e.g. aspirin). Certain medical disorders, such as liver and renal disease, are associated with a haemorrhagic state of multifactorial origin. The severity of the haemorrhagic diathesis is in general proportional to the severity of the underlying disorder. The maintenance of blood within the vascular system depends upon the integrity of the coagulation mechanism, the presence of a reasonable number of functional platelets and endothelium-lined vessels capable of constriction when severed.

Platelets are responsible for controlling the initial onset of haemorrhage by adhering to subendothelial components, for example collagen and microfibrils, and forming a plug in the severed vessel. Platelets circulate for 7–10 days after being released from bone marrow megakaryocytes. They have a complex structure suitable for responding rapidly to breaches in vascular integrity. Platelets have cell surface receptors for various activated components of the coagulation cascade (such as thrombin), and possess delta granules containing vasoactive amines (e.g. adenosine triphosphate and 5-hydroxytryptamine), and alpha granules containing protein components of the haemostatic system (e.g. von Willebrand factor and factor V). Inadequate platelet function or a low platelet count typically results in mucosal bleeding such as purpura, easy bruising, epistaxis, gastrointestinal haemorrhage or menorrhagia.

The von Willebrand factor is a plasma protein, secreted by endothelial cells, that promotes the adhesion of
platelets to damaged vessel walls. It also acts as a carrier protein for factor VIII; hence, in von Willebrand's disease, the plasma level of factor VIII is often reduced because, without its carrier, it is unstable and has a reduced plasma half-life. As the von Willebrand factor is essential for the adhesion of platelets to traumatized vessels, patients with von Willebrand's disease present with similar bleeding patterns to those of individuals with platelet functional disorders (i.e. mucosal haemorrhage).

The coagulation cascade consists of a series of proenzymes. Initially, each acts as an enzyme substrate; after activation, they act as enzymes activating the subsequent proenzyme in the cascade (Fig. B.6). The rate of many of the steps in the coagulation cascade is enhanced if these occur on the platelet surface, although these reactions can also happen in plasma. This procoagulant property is due to specific platelet receptors for components of the coagulation cascade.

Conventionally, the coagulation system is considered to be composed of two parts – the intrinsic and extrinsic components – although recent research has revealed that the system is considerably more complicated than is illustrated in Fig. B.6. Deficiencies in the coagulation system may be single (e.g. haemophilia) or multiple (e.g. warfarin therapy). Bleeding can occur due to the presence of an inhibitor (usually an IgG antibody) against one or more of the coagulation factors or platelets (e.g. in idiopathic thrombocytopenic purpura). Single coagulation deficiencies tend to cause haemarthrosis or muscle haematoma, while multiple abnormalities may cause almost any bleeding manifestation. In haemophilia, the primary haemostatic mechanism involving platelets is normal, so the bleeding often stops immediately after trauma; however, haemorrhage will then start several hours later because the platelet plug is not consolidated by the deposition of fibrin.

History-taking for a patient presenting with possible excessive bleeding should focus on the following important aspects:

- The duration of symptomatology indicates whether the possible haemorrhagic predisposition is congenital or acquired.
- The bleeding pattern indicates which component of the haemostatic system may be deficient: thrombocytopenia and platelet disorders give rise to purpura and bleeding into mucosal surfaces, while a coagulation defect usually results in muscle and joint haemorrhage.
- The timing of bleeding is also relevant. Bleeding starting at the time of trauma (e.g. dental extraction) indicates a failure of platelet plug formation, and therefore a platelet disorder or von Willebrand's disease.
- Spontaneous haemorrhage indicates a more severe bleeding disorder than that associated with provocation by trauma.
- Dental extractions, tonsillectomy and circumcision are all potent stressors of the haemostatic mechanism. If a patient has undergone any two of these procedures without excessive blood loss, they are unlikely to have a clinically significant bleeding problem.
- Family history is important, as many congenital conditions have a familial predisposition.
- The drug history is essential because almost all medicines can, by one mechanism or another, predispose towards bleeding. Ingestion of warfarin or aspirin has to be identified. Exposure to toxins or solvents at work or with hobbies may result in hypoplastic anaemia.
- The general medical history is essential to identify the many disorders that may result in thrombocytopenia or coagulation disturbances, such as liver disease, renal failure, connective tissue diseases and so on.

Clinical examination should identify all the sites of haemorrhage. The buccal cavity and optic fundi should therefore always be looked at, as superficial bleeding at these sites indicates severe platelet dysfunction. It may be necessary to use imaging procedures, for example computed tomography scanning or ultrasound, to fully document the extent of internal haematoma formation. Initial screening tests include a complete blood count, examination of a blood film, bleeding time, activated partial thromboplastin time (APTT; intrinsic system), prothrombin time (PT; extrinsic system), fibrinogen and D-dimers (a measure of fibrinolysis) (Table B.1).
If either the activated partial thromboplastin time (APTT) or the prothrombin time (PT) is found to be prolonged, the tests must be repeated after the addition of normal plasma, when the test time should become normal. Failure to normalize the clotting time raises the suspicion of the presence of an inhibitor.

Any patient with thrombocytopenia for which the cause is not immediately and unequivocally apparent should have a bone marrow aspirate and/or trephine performed. This will allow an assessment of megakaryocytic numbers. They are reduced in conditions of the underproduction of platelets (e.g. hypoplastic anaemia), but increased when there is increased destruction and/or pooling of circulating platelets (e.g. splenomegaly). A trephine biopsy is particularly useful for assessing whether the bone marrow is infiltrated with carcinoma cells (Table B.2).

### Table B.1 Haematological changes in bleeding-related conditions

<table>
<thead>
<tr>
<th>Condition</th>
<th>Platelet count</th>
<th>Bleeding time</th>
<th>APTT</th>
<th>Prothrombin ratio</th>
<th>Fibrinogen</th>
<th>D-dimer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thrombocytopenia</td>
<td>↓</td>
<td>↑</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>Von Willebrand's disease</td>
<td>N</td>
<td>↑</td>
<td>N or ↑</td>
<td>N</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>Haemophilia A or B</td>
<td>N</td>
<td>N</td>
<td>↑</td>
<td>N</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>Warfarin/liver disease</td>
<td>N</td>
<td>N</td>
<td>N</td>
<td>↑</td>
<td>N</td>
<td>N</td>
</tr>
<tr>
<td>Disseminated intravascular coagulation</td>
<td>↓</td>
<td>↑</td>
<td>↑↑</td>
<td>↑</td>
<td>↓</td>
<td>↑</td>
</tr>
</tbody>
</table>

### Table B.2 Causes of bleeding

<table>
<thead>
<tr>
<th>Cause</th>
<th>Example</th>
</tr>
</thead>
<tbody>
<tr>
<td>Structural lesion</td>
<td>Peptic ulcer, aortic aneurysm, arteriovenous malformation, subarachnoid haemorrhage, angiodysplasia (colonic bleeding), endometriosis, Osler–Weber–Rendu syndrome (hereditary haemorrhagic telangetasis – epistaxis or gastrointestinal bleeding)</td>
</tr>
<tr>
<td>Coagulation defect</td>
<td>Haemophilia, von Willebrand's disease, disseminated intravascular coagulation, thrombocytopenia (autoimmune, marrow failure)</td>
</tr>
<tr>
<td>Medication</td>
<td>Warfarin (especially with drug interactions), heparin (can also cause thrombocytopenia), aspirin</td>
</tr>
<tr>
<td>Systemic illness</td>
<td>Liver disease, renal failure, malabsorption (vitamin K deficiency)</td>
</tr>
<tr>
<td>Infection</td>
<td>Schistosomiasis (haematuria), tuberculosis (haematemesis), Ebola virus</td>
</tr>
<tr>
<td>Other</td>
<td>Snake bite venom, scurvy (vitamin C deficiency)</td>
</tr>
</tbody>
</table>

### BLOOD PRESSURE, HIGH

**Gerry Carr-White**

An isolated (casual) elevated blood pressure reading can have three possible explanations. These are: (i) an error due to either faulty or inappropriate apparatus or faulty technique (observer error); (ii) a temporary elevation of blood pressure at the time of measurement (elevation due to biological variability); and (iii) sustained blood pressure elevation in the individual not attributable to environmental stimuli (elevated basal pressure).

**OBSERVER ERROR**

This may be due to instrument design and maintenance, inadequate cuff size, the technique used for measurement, or the criteria used for determining systolic and diastolic pressure.

**Faulty or inappropriate apparatus**

Aneroid sphygmomanometers lose accuracy with time and require regular calibration. Dirt in the escape valve may cause irregular deflation, and add to any inaccuracy of reading. Incorrect readings may be obtained if, in the case of a mercury sphygmomanometer, the mercury column does not read zero before inflation. If the mercury column is not vertical, readings will overestimate blood pressure.

If the rubber bladder contained within the sphygmomanometer cuff is too short, the blood pressure will be overestimated as pressure is not fully transmitted to the artery. The bladder should therefore cover at least 80 per cent of the circumference of the arm. A 35 cm bladder is recommended for normal or lean arms, longer bladders (up to 42 cm) being necessary for heavily muscled or obese arms. Too narrow a bladder also leads to an overestimation of blood pressure, although this causes fewer problems than too short a bladder. The width of the bladder should be at least 40 per cent of the circumference of the arm.
Faulty technique
The cuff should be inflated to at least 30 mmHg above the point at which the radial pulse disappears. The cuff should then be deflated at a rate of 2 mmHg per second over the critical points. When using mercury sphygmomanometers, the eye should be level with the upper level meniscus, otherwise parallax will give rise to erroneous readings. Rapid re-inflation of the cuff or failure to deflate properly before repeating blood pressure measurement may increase the level at which the Korotkoff sounds appear, and so overestimate systolic blood pressure level. Rounding up or down to the nearest figure ending in a zero or five (digit preference) may make a small contribution to erroneous readings. Readings can normally be rounded to the nearest even number. A predetermined threshold for the diagnosis of hypertension or for treatment may also unconsciously influence the observer’s record (observer bias). The arm should be supported at the mid-sternal level. If the arm is held in a dependent position, diastolic and systolic blood pressures can be overestimated by up 10 mmHg.

Where the phase of muffling (phase IV Korotkoff sounds) is used for estimating the diastolic blood pressure level, values 5–10 mmHg higher are obtained compared with when the point of disappearance of the Korotkoff sounds is used (phase V Korotkoff sounds). Generally, phase V values correlate better with intra-arterial pressures, and reproducibility between observers is superior.

SUBJECT (BIOLOGICAL) VARIABILITY
Anxiety, recent physical activity, recent cigarette smoking, a cold temperature and physical pain all cause an elevation of blood pressure through activation of the autonomic nervous system. The first reading obtained by a doctor is usually higher than subsequent readings, either on the same occasion or on later occasions. Thus, significant blood pressure falls have been recorded with the passage of time in placebo-treated patients in clinical trials. These important pressor effects can be minimized by a careful explanation of the procedure to the patient beforehand, a comfortable environment, and allowing a 2- to 3-minute period of rest before the blood pressure is measured. The diagnosis of hypertension should not normally be made before blood pressure has been measured on three or more occasions, unless other evidence such as the presence of significant target organ damage is found, or unless very high blood pressure levels are observed. Some patients show a consistent pressor response to the presence of a doctor or to blood pressure recording (‘white coat’ hypertension).

This should be suspected where high blood pressure levels are repeatedly recorded in the absence of any fundal, electrocardiographic or echocardiographic evidence of hypertensive organ damage. It should also be suspected in patients who appear consistently tense or anxious during the measurement procedure. In these circumstances, ambulatory monitoring of blood pressure or self-monitoring at home using an electronic digital device should be used.

Blood pressures should, on the first occasion, always be measured in both arms, since minor degrees of inequality are quite common. If there is a reproducible difference of 20 mmHg for systolic blood pressure and 10 mmHg for diastolic blood pressure, simultaneous measurements should be carried out: the higher values should be taken as more representative for the patient’s clinical management. The time at which antihypertensive drugs are taken may also influence blood pressure. For patients receiving once-daily treatment, it is probably best to measure blood pressure just before the patient takes the daily dose.

RAISED BLOOD PRESSURE – ASSESSMENT
In unselected populations, blood pressure is distributed as a smooth unimodal curve. There is therefore no natural line of demarcation between normal and abnormal blood pressures in unselected subjects. The incidence of cardiovascular disease (i.e. stroke, ischaemic heart disease and peripheral vascular disease) is related to blood pressure level in a curvilinear fashion with no evidence of a lower threshold down to a systolic blood pressure of approximately 115 mmHg and a diastolic blood pressure of around 75 mmHg. It is therefore impossible to define hypertension by reference to a value above which a patient is at risk. The level of blood pressure at which drug treatment is indicated varies somewhat from country to country, although clinicians are aided by national guidelines.

There are nevertheless great clinical advantages in selecting (albeit arbitrary) thresholds for a diagnosis of hypertension. The most commonly used criteria of recent times are those of the World Health Organization (Vth Korotkoff phase). These are:

- Normal range: equal to or below 140/90 mmHg
- Hypertensive range: 160/95 mmHg and above
- Borderline or intermittent: 140–159/90–94 mmHg

More recently the European Society of Cardiology states:

Hypertension is defined as systolic BP ≥140 mmHg and/or diastolic BP ≥90 mmHg. The diagnosis of
hypertension should be based on at least two BP measurements in the sitting position per visit on at least two visits.

Cut-offs for the definition of hypertension by home monitoring (and in particular by 24-hour BP monitors) are: 130/80 mmHg for 24-h BP, 135/85 mmHg for daytime ambulatory BP and home BP, and 120/70 mmHg for night-time BP.

The presence of systodiastolic hypertension, diastolic hypertension or (isolated) systolic hypertension is associated in all cases with increased cardiovascular risk in individuals, and the decision to treat is nowadays made with reference to national guidelines based upon an assessment of absolute cardiovascular risk.

Since both systolic and diastolic blood pressures rise with age (Fig. B.7), the apparent prevalence of hypertension will also rise with age. Furthermore, where a single reading is used, rather than average reading over several measurements, the apparent prevalence of hypertension will be much higher, since blood pressure tends to fall with repeated measurements (see above). Thus, approximately 40 per cent of untreated middle-aged and elderly men will have a diastolic blood pressure of 90 mmHg or over on single readings, but this figure will fall to 15–20 per cent on repeated measurement.

There is little justification in taking age into account in defining hypertension in the adult population since the risks of high blood pressure, at least up to extreme old age (i.e. 80 years and above), are related to absolute blood pressure level. In children, however, the same criteria used to define hypertension in adults clearly cannot be applied. Blood pressure rises rapidly in the first few days of life, and slightly more slowly over the next few weeks, and then shows little change between 6 weeks and 6 years of age, when blood pressure begins to rise again slowly. Thus, in infancy, blood pressures of 80–90/50–60 mmHg are observed, and in children aged 10, blood pressures of 90–100/60–70 mmHg.

Blood pressure levels that would be acceptable in adults are poorly tolerated by children, and the arbitrary values used in adults therefore have to be adjusted and related to age. The American Task Force on Blood Pressure Control in Children has recommended that sustained blood pressure levels (obtained on at least three separate occasions) above the 95th centile for age should be considered abnormal in this context.

The malignant or accelerated phase may occur in hypertension from any cause. Although the term ‘malignant hypertension’ was previously reserved for patients with papilloedema, it is now recognized that, when untreated, the prognosis is just as bad in patients who show hypertensive haemorrhages or exudates (grade III retinopathy), and it is now customary to term hypertension associated with grade III or IV retinopathy as malignant. Malignant hypertension is more common in males. It is usually associated with a diastolic blood pressure of 130 mmHg or more and, compared with other hypertensive patients, these patients are relatively young (most usually 40–50 years of age). The clinical picture of malignant hypertension reflects the pathological process, i.e. acute severe vascular damage. Symptoms are much more frequent than with ‘benign’ essential hypertension. They include blurring of vision, mental impairment, haematuria or haematospermia, and clinical features of target organ damage.

Hypertensive haemorrhages in malignant hypertension are of two sorts. Flame-shaped haemorrhages are more superficial and owe their character to constraints imposed on nerve fibres. Dot and blot haemorrhages are deep to the nerve fibres, so are not limited in the same way. Haemorrhages are a sign of recent severe vascular damage, and usually disappear after a few weeks of effective blood pressure control. Exudates are of two types. Hard or waxy exudates represent the end result of fluid leakage into the fibre layers of the retina from damaged vessels, often with associated nerve fibre damage. Fluid is reabsorbed, leaving a protein lipid residue that is slowly removed by macrophages, finally leaving a hyaline deposit that may sometimes persist. Like retinal haemorrhages, hard exudates are forced into a radial spoke-like distribution by the nerve fibres around the macula (‘macular star’). Soft exudates or cotton-wool patches are quite different aetio logically and ophthalmoscopically. They are usually larger than hard exudates and have a woolly,
ill-defined edge. They are not true exudates but nerve fibre infarcts caused by hypertensive vascular occlusion. Unlike hard exudates, these lesions disappear within a few weeks of establishing adequate hypertensive therapy. Papilloedema is associated with increased pressure within the optic disc secondary to severe vascular damage. Venous distension is followed by increased vascularity of the optic disc, which has a pink appearance, with blurring of the disc margins and loss of the optic cup. Raising of the optic disc with anterior displacement of the vessels occurs later. Often the surrounding retina shows oedema, small radial haemorrhages and cotton-wool exudates.

RAISED BLOOD PRESSURE – CAUSES

Essential hypertension

No identifiable cause can be found in the vast majority of patients who present with sustained blood pressure elevation (essential hypertension). High blood pressure in such individuals is believed to be multifactorial, with a substantial contribution from both genetic and environmental factors. Studies of the prevalence of hypertension in families and in monozygotic and dizygotic twins have confirmed the importance of genetic factors. The advent of the ‘molecular medicine’ era has heralded the development of newer techniques to study mechanisms and genes determining blood pressure control. To date, genome-wide screening has identified several promising chromosomal regions that will help to offer mechanistic insights into how blood pressure is controlled and that may give rise to novel therapeutic targets for antihypertensive drugs.

Epidemiological studies have also emphasized the importance of other factors that are associated with blood pressure elevation. These include dietary salt, obesity and heavy alcohol intake. It appears that all ethnic groups are susceptible to hypertension, but that the expression of frank cardiovascular disease differs between, for instance, African-Caribbean individuals (who have a higher risk of stroke) and South Asian individuals (who have a higher incidence of coronary heart disease). An extensive investigation of patients with essential hypertension may show a variety of minor changes, for example a slight elevation in hematocrit, an increase in renal vascular resistance, decreased renal plasma flow and an increased filtration fraction. All these changes are, however, probably secondary to structural changes induced in the blood vessels by hypertension. One possible clue to the aetiology is provided by evidence of sympathetic nervous systemic activation in young hypertensives, who often have a cardiac output in the upper part of the normal range, an increased heart rate and slightly elevated circulating noradrenaline (norepinephrine) levels. Later in the course of hypertension, this evidence for increased nervous system activity disappears, and blood pressure is maintained by elevated peripheral resistance alone. There is no evidence that other powerful pressor systems, such as the renin–angiotensin system, or sodium retention are responsible for blood pressure elevation in essential hypertension.

The vast majority of patients with essential hypertension are diagnosed either at routine examination, or incidentally when attending a doctor for other medical problems. The early stages of hypertension are in most cases asymptomatic. Occipital headaches that are throbbing in nature and worse in the morning are described as classical but are only seen in a small minority of patients. In most cases, headaches are probably unrelated to hypertension, although the frequency of headaches in a hypertension clinic decreases with effective blood pressure control. Epistaxes are more frequent in hypertensive patients, but they are an unusual manifestation. The other manifestations of hypertension are due to target organ damage. Dyspnoea due to left ventricular failure is a very late manifestation. Besides reflecting an increased load against which the left ventricle is working, it may also be due to associated ischaemic heart disease, which is of course more common in hypertension. Visual disturbances are only seen with advanced (grade III or grade IV) retinopathy, although, occasionally, arteriovenous nipping, seen in less severe retinopathy, can cause a branch retinal vein occlusion. Renal impairment is uncommon in patients who have only grade I or II retinopathy. Nocturia is, however, frequently seen in hypertension of all grades, reflecting a disturbance in the normal circadian rhythm of urine formation. Focal neurological signs may either reflect a cerebrovascular accident (cerebral haemorrhage or thrombosis) or be due to focal oedema (hypertensive encephalopathy). This disorder is characterized by transient focal neurological signs associated with very high blood pressure levels. It is due to systemic blood pressure exceeding the upper autoregulatory range of cerebral blood flow control, so that focal hyperaemia and oedema occur. Peripheral vascular disease is due to hypertension-induced atheroma in the large arteries and aorta.

Secondary hypertension

High blood pressure can be attributed to a specific disorder or drug only in a minority of patients. The quoted incidence of secondary hypertension has
renal artery or its smaller branches (renovascular cause renal hypertension. These are diseases of the renal parenchyma, and renin-secreting tumours, which are derived from the cells of the juxtaglomerular apparatus. The last named is an extremely rare cause of hypertension occurring in children or young adults. There are also case reports of hypersecretion of renin by a Wilms’ tumour, renal carcinoma, bronchial carcinoma and pancreatic adenocarcinoma. Hypertension is caused by the high renin levels, with associated secondary aldosteronism, although structural changes in the resistance vessels help to maintain blood pressure when hypertension has been maintained for prolonged periods. Renovascular and renoparenchymal hypertension are not entirely discrete categories. Thus, renal parenchymal disease such as pyelonephritis or glomerulonephritis gives rise to renal ischaemia, and hypersecretion of renin can frequently be demonstrated. The other known factor that plays a role in some patients with bilateral renovascular or renoparenchymal disease (or disease in a single kidney) is sodium retention. This is particularly notable in acute glomerulonephritis and in advanced renal failure, where oedema is often associated with hypertension. Unfortunately, from the diagnostic point of view, in many patients with renovascular or renoparenchymal disease, there are neither high renin levels nor evidence of sodium retention. In some cases, it seems likely that chronic hypertension has given rise to structural changes in the resistance vessels, which then maintain blood pressure even after the precipitating factor is no longer in evidence. It also seems likely, however, that the kidney regulates blood pressure in other less well understood ways. For instance, the renal medulla secretes vasodepressor material, and this mechanism may be impaired in some forms of renal hypertension.

Because of the multiplicity of renal mechanisms, and because secondary changes may maintain blood pressure even after the initial mechanism has ceased to act, the diagnosis of renal hypertension is frequently extremely difficult. Clinical and biochemical features are often conspicuous by their absence. Certain clues may, however, be suggestive. Thus, severe hypertension presenting in a young patient (e.g. below the age of 30) in the absence of a family history of hypertension makes a renal cause more likely. A renal cause is more likely to be found in patients with malignant hypertension, and in patients whose blood pressure rises rapidly. A renal bruit, particularly when it occurs both in the diastolic and systolic phases, is more suggestive of a renovascular cause, although such bruits are frequently heard in the absence of hypertension, diseases of the renal parenchyma, and renin-secreting tumours, which are derived from the cells of the juxtaglomerular apparatus. The last named is an extremely rare cause of hypertension occurring in children or young adults. There are also case reports of hypersecretion of renin by a Wilms’ tumour, renal carcinoma, bronchial carcinoma and pancreatic adenocarcinoma. Hypertension is caused by the high renin levels, with associated secondary aldosteronism, although structural changes in the resistance vessels help to maintain blood pressure when hypertension has been maintained for prolonged periods. Renovascular and renoparenchymal hypertension are not entirely discrete categories. Thus, renal parenchymal disease such as pyelonephritis or glomerulonephritis gives rise to renal ischaemia, and hypersecretion of renin can frequently be demonstrated. The other known factor that plays a role in some patients with bilateral renovascular or renoparenchymal disease (or disease in a single kidney) is sodium retention. This is particularly notable in acute glomerulonephritis and in advanced renal failure, where oedema is often associated with hypertension. Unfortunately, from the diagnostic point of view, in many patients with renovascular or renoparenchymal disease, there are neither high renin levels nor evidence of sodium retention. In some cases, it seems likely that chronic hypertension has given rise to structural changes in the resistance vessels, which then maintain blood pressure even after the precipitating factor is no longer in evidence. It also seems likely, however, that the kidney regulates blood pressure in other less well understood ways. For instance, the renal medulla secretes vasodepressor material, and this mechanism may be impaired in some forms of renal hypertension.

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any lesions of the renal arteries. Generalized oedema suggests acute glomerulonephritis. This may be post-streptococcal, or may be a manifestation of systemic disease such as Henoch–Schönlein purpura, Goodpasture's syndrome, polyarteritis nodosa or Wegener's granulomatosis. Clinical evidence of uraemia, perhaps associated with dependent oedema, suggests advanced renal disease, most probably due to end-stage chronic glomerulonephritis or chronic pyelonephritis. It has to be borne in mind, however, that severe hypertension can give rise to hypertensive nephropathy and renal failure, so that uraemia can be an effect rather than a cause of hypertension. This is more likely in patients with grade III or IV retinopathy. Although a history of urinary tract infections, perhaps in childhood or many years previously, may suggest chronic pyelonephritis, in the majority of cases of chronic pyelonephritis and hypertension there is no previous history of urinary tract infection. In some patients, there may be a history of previous reflux uropathy in childhood. A history of renal disease in the family suggests polycystic kidneys or, less commonly, hereditary nephritis. Relevant features in the history may less commonly indicate such causes as gouty nephropathy, diabetes, irradiation nephritis, amyloidosis, renal tuberculosis or heavy metal poisoning.

In only a minority of cases with renal hypertension will the history and examination yield the diagnosis. Urine dipstick testing and microscopy are simple, useful and inexpensive investigations that may provide evidence of renal pathology. Measurement of serum electrolytes, urea and creatinine may indicate the presence of renal disease in a few patients. Plasma renin and aldosterone are often normal, and may indeed be subnormal where renin secretion has been suppressed by sodium retention. In addition, severe and malignant hypertension are often associated with elevated plasma renin levels even where there is no primary renal disease. Further investigation of renal hypertension requires renal imaging with ultrasonography in the first instance. Screening for renovascular hypertension involves duplex ultrasound, nuclear renal imaging, magnetic resonance angiography or computed tomographic (CT) angiography, depending upon which centre the patient is in. Definitive diagnosis of the lesion in renovascular hypertension demands renal angiography. There is considerable debate among medical professionals as to how best to manage renovascular hypertension, although more recent evidence seems to suggest that renal artery stenting is of no benefit in atherosclerotic renovascular disease.

Hypertension in pregnant women
Hypertensive disorders are the most common medical complications of pregnancy, and are broadly speaking classified into three varieties: chronic hypertension, gestational hypertension and pre-eclampsia. Pre-eclampsia is unique in being a self-limiting form of hypertension. Classical pre-eclampsia occurs for the first time during the third trimester of pregnancy, and blood pressure falls immediately after delivery. Blood pressure elevation is associated with significant proteinuria and oedema. Classically, pre-eclampsia is observed during the first pregnancy only, and is more common in older women and those with diabetes, multiple pregnancies or a hydatidiform mole. It has to be differentiated from essential hypertension that has been exacerbated by pregnancy (chronic hypertension). In this case, hypertension occurs during the first trimester, and becomes progressively worse with successive pregnancies. The difference between the two conditions is often not clear-cut, and diagnosis may have to be delayed until the course of blood pressure after gestation can be observed. Gestational hypertension is defined as the development of hypertension without features of pre-eclampsia after 20 weeks’ gestation in a previously normotensive woman. This form of hypertension tends to be associated with a good prognosis, and in many instances may not even require blood pressure-lowering therapy.

Endocrine and metabolic hypertension
Primary aldosteronism is associated with either a single adenoma or bilateral hyperplasia of the adrenocortical zona glomerulosa. The latter may take the form of diffuse hypertrophy, or there may be multiple small adenomata (micronodular hyperplasia). It is important to distinguish between a single adenoma and hyperplasia as the treatment of the first is surgical, and the second medical. Very rarely, the lesion may be carcinomatous, and occasional cases have been described with no histological lesion. Primary aldosteronism probably accounts for less than 1 per cent of cases of hypertension. The clinical picture may be indistinguishable from essential hypertension. The most characteristic symptom is generalized muscle weakness, although cramps, tetany and polyuria occasionally occur. Malignant hypertension is comparatively rare in primary aldosteronism, perhaps because the rise in blood pressure is gradual rather than rapid. The biochemical features that suggest primary aldosteronism are a low serum potassium associated with serum sodium that is in the upper part of, or just above, the normal range. Further investigations will demonstrate a suppression
of plasma renin and an elevation of blood and urinary aldosterone. It is important to differentiate primary and secondary aldosteronism (which is frequently seen in severe hypertension or in diuretic-treated patients). In secondary aldosteronism, the serum sodium is low, and the plasma renin elevated. Suprarenal tumours are currently best visualized by contrast-enhanced CT scanning, and differential suprarenal venous sampling can then be undertaken to determine whether the hyperaldosteronism is unilateral or bilateral.

Very rarely, cases have been described with an isolated secretion of other mineralocorticoids, such as deoxycorticosterone. 11-beta-hydroxylase deficiency occurs in children and is associated with virilization (adrenogenital syndrome). 17-alpha-hydroxylase deficiency is associated with sexual immaturity as the production of sex hormones is impaired. In all these conditions, mineralocorticoid-induced sodium retention and hypokalaemia occur, with a suppression of plasma renin. Liddle’s syndrome is an autosomal dominant condition characterized by early-onset hypertension with hypokalaemic alkalosis. It is caused by activating mutations in the epithelial sodium channels in the renal collecting tubules, leading to increased sodium-retaining and potassium-secreting activity. Thus, although the biochemical features of primary aldosteronism are present, with a low potassium and renin level, aldosterone is also very low. Glucocorticoid-remediable hyperaldosteronism (also called dexamethasone-suppressible hyperaldosteronism) is another autosomal dominant trait in which early-onset hypertension is associated with haemorrhagic stroke. It has the biochemical features of primary aldosteronism, but the biochemistry is corrected by suppressing adrenocorticotrophic hormone (ACTH) with dexamethasone. This is due to a mutation producing a chimeric gene encoding a protein with aldosterone synthase enzymatic activity whose expression is regulated by ACTH. Other rare causes of heritable hypertension include the syndrome of apparent mineralocorticoid excess (autosomal recessive) and type II pseudohypoaldosteronism (Gordon’s syndrome, autosomal dominant).

The hypertension of Cushing’s syndrome is usually associated with a hypersecretion of glucocorticoids, although mineralocorticoids such as deoxycorticosterone may also occasionally be elevated. The hypertension arises due to the cortisol-mediated activation of the mineralocorticoid receptor secondary to supersaturation of the enzyme 11-beta-hydroxysteroid dehydrogenase type 2, which would normally inactivate cortisol by metabolizing it to cortisone. Phaeochromocytomas are tumours of sympathetic tissue that produce hypertension by the secretion of catecholamines (adrenaline, noradrenaline and, occasionally, dopamine). They are probably responsible for hypertension in less than 0.1 per cent of hypertensive patients. Although the tumours usually originate in the suprarenal medulla (when adrenaline secretion tends to predominate), they may also arise from sympathetic ganglia associated with the abdominal, and rarely thoracic, aorta and the bladder wall. Tumours are frequently multiple and occasionally malignant. Classically, suspicion of phaeochromocytoma is raised on clinical grounds. The patient has short periods of high blood pressure associated with other features of sympathetic activity. These include sweating, flushing and a throbbing headache, abdominal or chest pain and weight loss. The anxiety usually associated with such attacks is presumably a manifestation of visceral feedback. Examination during attacks usually shows either tachycardia or bradycardia, and severe hypertension. Attacks may be provoked by specific movement, exercise, micturition, abdominal palpation or surgery (Fig. B.8); indeed, occasional fatalities have been described during clinical examination. In addition, some patients have a low standing blood pressure, and this may cause symptoms of postural hypotension. Occasionally, very high circulating catecholamine levels have been associated with a condition that resembles cardiogenic shock, and post-mortem focal myocardial lesions have been observed. Glucose tolerance is frequently impaired, basal metabolic rate is elevated and free fatty acid levels are reduced.

**Figure B.8** Systolic blood pressure fluctuation during manipulation of the tumour at operation in a patient with a phaeochromocytoma.
raised. In some cases, the tumour is associated with neurofibromatosis (von Recklinghausen's disease). A rare combination of endocrine disorders is inherited as an autosomal dominant; this comprises multiple phaeochromocytomas, medullary carcinoma of the thyroid and hyperparathyroidism (Sipple's syndrome or multiple endocrine adenomatosis type II). Other associations are tuberose sclerosis and Sturge–Weber syndrome. Diagnosis is made by finding significantly elevated catecholamines or catecholamine metabolites (such as vanillylmandelic acid or normetadrenaline and metadrenaline) in the urine. Significant numbers of false-negative results occur, particularly if urine collection does not coincide with a pressor attack. For this reason, if the diagnosis is seriously being considered, multiple urine collections are necessary. Additionally, plasma catecholamines can be measured while the patient is under observation. Elevated levels associated with tachycardia and raised blood pressure are suggestive. The main differential diagnosis here is anxiety. The administration of clonidine results in a lowering of plasma catecholamine levels in patients with anxiety, but has no effect upon the high catecholamine levels observed in phaeochromocytoma. Tumours can be imaged either by CT scanning or by isotope studies using metaiodobenzguanidine. If no suprarenal tumour is seen, venous sampling for catecholamines at different levels in the inferior and superior vena cava may help to locate an ectopic tumour.

Blood pressure is elevated in about 30 per cent of patients with acromegaly, although different clinical series quote a wide-ranging prevalence of 20–60 per cent. The exact mechanisms underlying the development of hypertension in acromegaly remain unclear, although it is recognized that growth hormone induces sodium retention. Consistent with this finding is the observation that plasma renin is often suppressed in acromegalic patients. There is also an association between suprarenal adenomata and acromegaly, which may be relevant in some patients. High blood pressure is approximately twice as common in hypothyroid patients as in the general population. In most cases, blood pressure falls when thyroid deficiency is corrected. Hypertension cannot be clearly related either to increased renin secretion or to sodium and water retention. It has been suggested that there may be an abnormality in vascular smooth muscle produced by thyroid hormone deficiency.

Other causes of hypertension

Coarctation of the aorta is the most common cardiovascular cause of hypertension. In coarctation of the aorta, there is narrowing, usually situated distal to the origin of the left subclavian artery at or near the insertion of the ligamentum arteriosum (Fig. B.9). The so-called infantile type of coarctation in which the ductus is patent is irrelevant to hypertension. Uncomplicated coarctation may present as mild blood pressure elevation in childhood, and occasionally hypertension becomes suddenly more severe. Infants with other cardiac lesions present with congestive failure. The young adult usually presents with asymptomatic hypertension or with the complications of hypertension. Unless there is an associated cardiac lesion, the clinical symptoms are usually indistinguishable from those of essential hypertension beginning at an unusual age, although diminished circulation through the legs may cause cramps. Physical signs that suggest the diagnosis are raised blood pressure in the arms associated with a normal or low blood pressure in the legs (a wide-leg cuff has to be used to determine this), delayed, weak or absent femoral pulses, an ejection systolic murmur heard best posteriorly between the left scapula and the spine, and pulsatile collateral vessels situated around the scapulae and in the posterior intercostal spaces – these are more noticeable on sitting the patient forwards. There are usually associated bruits. There may also be an aortic systolic ejection murmur due to a bicuspid aortic valve, which is present in 50 per cent of cases. A firm diagnosis can often be made from the chest X-ray. The characteristic double aortic knuckle is made up of the dilated left subclavian artery and post-stenotic
dilatation of the descending aorta. Another almost pathognomonic sign is notching of the lower borders of the ribs. These are not to be confused with defects of an erosive nature in the superior margins of the ribs, seen rarely in poliomyelitis, hyperparathyroidism, rheumatoid arthritis and scleroderma. The diagnosis can finally be confirmed, and the extent of the coarctation delineated, by CT aortography.

Neurological disease only very rarely causes hypertension. Raised intracranial pressure probably causes hypertension through brainstem compression and ischaemia, activating sympathetic efferent outflow from the vasomotor centre. Transient blood pressure elevation may be seen after head injury, presumably for the same reason. Vascular disease, brainstem encephalitis and poliomyelitis also occasionally produce hypertension through involvement of the brainstem centre. Lesions of the upper part of the spinal cord may cause severe hypertension through interference with cardiovascular reflexes. Such hypertension may be paroxysmal, as a result of an acute pressor response to stimulation of viscera such as the bladder or rectum.

In addition to the contraceptive pill and corticosteroid therapy, many drugs can either cause hypertension or exacerbate pre-existing hypertension. Hypertension and tachycardia are common findings in patients attending Accident and Emergency Departments who have abused recreational drugs such as ecstasy and cocaine. Non-steroidal anti-inflammatory drugs raise blood pressure probably through an inhibition of renal prostaglandin synthesis, which plays a role in the regulation of sodium and water output. Usually the degree of blood pressure elevation is mild, but the administration of non-steroidal anti-inflammatory drugs may cause a loss of blood pressure control in patients on antihypertensive therapy. Antimigraine preparations of the serotonergic variety (5-hydroxytryptamine type 1 agonists or ‘triptans’) can produce transient increase in blood pressure.

Nowadays rarely used, monoamine oxidase inhibitors such as phenelzine and tranylcypromine can produce severe paroxysmal hypertension. This is particularly likely to occur when used in combination with tyramine-containing foods (of which mature cheese is the most important) and amphetamines. Clinically, the resulting syndrome resembles that observed in phaeochromocytoma. Sympathomimetic amines (e.g. amphetamine, ephedrine, metaraminol and other synthetic agents) are often used as nasal decongestants and may cause significant hypertension. Liquorice and carbenoxolone raise blood pressure through inducing a syndrome that resembles primary aldosteronism with a hypokalaemic alkalosis; in addition, the patients may be oedematous. This is due to direct inhibition of 11-beta-hydroxysteroid dehydrogenase type II, which normally metabolizes cortisol to cortisone and thereby allows the mineralocorticoid effects of cortisol (which has equal affinity to aldosterone for the mineralocorticoid receptor) to predominate. Ciclosporin and erythropoietin cause high blood pressure by mechanisms that have not been elucidated. Withdrawal of the centrally acting anti hypertensive agent clonidine causes paroxysmal hypertension due to increased efferent sympathetic nerve activity. Certain medical preparations contain large amounts of sodium. These include resonium A, para-aminosalicylate, sodium carbenicillin and some antacid mixtures. Whereas these have no effect on blood pressure in healthy individuals, hypertension may be caused in patients with renal failure.

**BLOOD PRESSURE, LOW**

Mark Kinirons

The lower half of the blood pressure distribution curve in healthy unselected populations is just as smooth as the upper half with no evidence for a discrete group of hypotensive subjects (see BLOOD PRESSURE, HIGH, p. 58). The diagnosis of low blood pressure is therefore just as arbitrary as the diagnosis of high blood pressure. The clinical significance of the diagnosis is, however, quite different. Both high and low blood pressure are most commonly multifactorial, i.e. the result of the interaction of genetic and environmental factors (biological variability). High blood pressure carries an increased cardiovascular risk although it is only infrequently due to a specific disease. Low blood pressure, however, when it is attributable only to biological variability, carries a lower risk of cardiovascular disease than the population average. Its only clinical significance therefore is when it causes symptoms (see FAINTS, p. 195) or when it is a manifestation of disease. In epidemiological studies it has been associated with increased prevalence of psychoneurotic symptoms. Additionally, in some studies, low diastolic blood pressure in treated hypertensive patients has been associated with an increased risk of cardiac death. Whether this is a result of treatment or not is controversial. The causes of low blood pressure are listed in Box B.5.

Low blood pressure can result from underactivity of any of the systems which maintain blood pressure. Since these systems assume particular importance when the subject stands, postural hypotension may
Cardiac
- Disturbances of rate and rhythm
  - Heart block
  - Dysrhythmias
- Obstruction to flow
  - Aortic or pulmonary valvular stenosis
  - Hypertrophic obstructive cardiomyopathy
  - Atrial myxoma
  - Primary pulmonary hypertension
  - Pulmonary embolism
  - Cardiac tamponade
- Mitral and tricuspid stenosis
- Cor triatrum
- Tetralogy of Fallot
- Eisenmenger’s syndrome
- Impaired ventricular function
  - Myocardial infarction
  - Cardiomyopathy

Impaired vasomotor control
- Vasovagal syncope
- Glossopharyngeal neuralgia
- Micturition, deglutition or post-tussive syncope
- Baroreceptor dysfunction in the elderly
- Autonomic degeneration (diabetes and Shy–Drager syndrome)
- Carotid sinus hypersensitivity

Impaired venous return
- Haemorrhage and dehydration
- Muscle wasting and prolonged bed rest

Metabolic and endocrine
- Phaeochromocytoma
- Serotonin-secreting tumours
- Hyporeninaemic hypoaldosteronism

Drugs
- Anti-hypertensives (particularly centrally acting agents, ganglion blockers, post-adrenergic ganglion blockers, alpha blockers and diuretics)
- CNS depressants
- Quinidine and other cardiac depressant drugs

be the only manifestation of low blood pressure. It is seen commonly, therefore, when fluid is lost from the gastrointestinal tract as a result of vomiting or diarrhoea, when renal fluid losses occur as in the excessive use of diuretics, Addison’s disease or in some patients with chronic pylonephritis and a sodium-losing tendency. It is observed when fluid is lost as a result of bleeding or burns. In wasting conditions or after prolonged bed rest, venous return to the heart and cardiac output may be reduced as a result of loss of skeletal muscle bulk; a low blood pressure is therefore frequently seen in this situation. Hypotension may result less commonly when cardiac output is reduced as a result of primary cardiac disease or cardiac tamponade, or as a result of obstruction to outflow of blood from the right or left side of the heart from, for example, valvular lesions.

Impairment of autonomic circulatory reflexes is often observed in diabetics and elderly patients with postural hypotension. In the latter case this is probably due to rigidity of the carotid artery and aorta in the region of the baroreceptors. Lesions of central pathways less commonly cause hypotension. Efferent pathways are interfered with particularly by ganglion blocking drugs and alpha-blocking agents such as prazosin or teroxzin. Degeneration of sympathetic pathways occurs in the rare Shy–Drager syndrome. The renin–angiotensin system does not assume great importance in blood pressure control unless patients are fluid depleted, so inhibition of this system does not normally cause a low blood pressure. However, a syndrome of hyporeninaemic hypoaldosteronism has been described in elderly subjects with postural hypotension.

**BODIES IMAGE, DISORDERS OF**
Andrew Hodgkiss

Broadly speaking, disorders of body image can be divided into two categories, differentiated by whether or not the problem is the presenting complaint for which the patient seeks correction (Box B.6).

**NOT THE PRESENTING COMPLAINT**
The classical organic presentation is hemiasomatognosia, the unilateral misperception of one's own body that is associated with parietal lobe lesions. A conscious form of hemiasomatognosia is the unilateral misperception of the patient believes and behaves as if half the body no longer exists. This can be subdivided into three forms:

**Box B.6 Causes of disorders of body image**

<table>
<thead>
<tr>
<th>Not the presenting complaint</th>
<th>The presenting complaint</th>
</tr>
</thead>
<tbody>
<tr>
<td>Most common</td>
<td>Common</td>
</tr>
<tr>
<td>Anorexia nervosa</td>
<td>Body dysmorphic disorder</td>
</tr>
<tr>
<td>Bulimia nervosa</td>
<td>Stress reaction</td>
</tr>
<tr>
<td>Parietal lobe lesions</td>
<td>Rare</td>
</tr>
<tr>
<td>Temporal lobe epilepsy</td>
<td>Gender dysphoria</td>
</tr>
<tr>
<td>Drug-related</td>
<td>Schizophrenia (with</td>
</tr>
<tr>
<td>Migraine</td>
<td>dysmorphophbic delusions)</td>
</tr>
<tr>
<td>Depersonalization</td>
<td>Monosymptomatic</td>
</tr>
<tr>
<td>Schizophrenia</td>
<td>hypochondriacal psychosis</td>
</tr>
<tr>
<td>LSD</td>
<td>disorders)</td>
</tr>
<tr>
<td>Mescaline</td>
<td>Psychotic depression</td>
</tr>
</tbody>
</table>
BODY IMAGE, DISORDERS OF

- Anosognosia for left hemiplegia, the denial of the existence of paralysis, which is usually confined to the first 2 weeks after onset (right parietal lobe lesion)
- Neglect syndromes, hemi-inattention and spatial neglect that is permanent (lesion of either parietal lobe)
- Gerstmann’s syndrome, which consists of finger agnosia, acalculia, agraphia and right/left disorientation, and is associated with autotopagnosia, the failure to localize, recognize or name parts of the body (left parietal lobe lesion)

Macro- or microsomatognosia, the experience that parts (or the whole) of the body have enlarged or shrunk, is associated most notably with temporal lobe epilepsy, but may also occur in depersonalization, migraine, schizophrenia and LSD or mescaline abuse. Similarly, changes in body shape, weight, colour or familiarity may occur in any of these conditions.

Overestimation of body width is a recognized feature of both anorexia nervosa and bulimia nervosa. The patient’s overestimation of size does not involve their height or other people’s body widths. It is an inconsistent finding among patients, although characteristic of the group, while its clinical significance lies in observations that it is a useful indicator of treatment response and the likelihood of relapse. This phenomenon has been reported to a lesser extent in young women without eating disorders, and it is postulated that overestimation of body width reflects the degree of personal value attached to body shape and size.

THE PRESENTING COMPLAINT

In gender dysphoria (or transsexualism), the individual’s dissatisfaction with their body stems from a deeply held belief that they belong to the opposite gender from their physical sexual characteristics. Sex reassignment surgery is sought, so that the external sexual characteristics of the desired sex may be acquired. These individuals usually live, dress and act as if they belong to their chosen gender, and report feeling relaxed and at peace, rather than sexually aroused, when doing so. Prolonged specialist assessment is always required before proceeding to definitive surgery.

A more common presenting complaint of body image disorder is body dysmorphic disorder (or dysmorphophobia), in which the patient asserts to being physically misshapen or defective in some way – which cannot be substantiated or is grossly exaggerated upon objective examination – and seeks cosmetic surgery. An important aspect of this condition is the concern, sometimes the certainty, that the defect is noticed by others. The most frequent sites for complaint are the nose or breasts, but the ears, chin, other facial features and genitals are not uncommon. While recognizing that a dysmorphophobic presentation is usually straightforward, it can be difficult – and sometimes impossible – to determine whether the belief is held with delusional conviction or is an overvalued idea. Generally, the more unusual the site and the more bizarre the belief, the more likely the presentation is a delusion, occasionally in the setting of psychotic depression, but more often as a feature of schizophrenia or monosymptomatic hypochondriacal delusional state. When doubt exists about whether the presentation is delusional, a treatment trial with a neuroleptic has been advocated and reported to be effective, although many patients will baulk at the prospect of this management approach.

The second problem is to determine whether cosmetic surgery will help the patient whose dysmorphophobia is neurotically or personality-based. It used to be considered that surgical correction in dysmorphophobic patients was inappropriate as it would neither affect their beliefs, nor prevent the psychological problems that many of them subsequently suffered. However, it has become apparent, that, following corrective surgery for minor deficits, the level of psychological disturbance falls, and the change in appearance is frequently regarded as satisfactory by the patient. This has led to an important distinction developing between patients who have trivial deformities and those who have none, but such a delineation can prove difficult to define.

In patients who have minor defects, the key is to understand the significance of the problem from the patient’s perspective. A middle-aged, male doctor is likely to view a small bump on the nose quite differently from a teenage girl, and it is important to acknowledge that it is her viewpoint that should be the more relevant. Sometimes minor physical defects can have major cultural, social or financial implications for the patient. Patients who should not be operated upon are, first, of course, those in whom corrective surgery is technically inappropriate either because there is no defect, or because the likelihood is that surgery will make matters worse or unchanged, or there is no corrective procedure for the patient’s complaint. Second, patients who have vague complaints and no specific treatment in mind, yet expect surgery both to be performed and to create perfection, should not be operated upon; most of these individuals suffer fundamentally from personality disorders – oversensitive, insecure, schizoid, narcissistic and obsessional traits have all been described. Third, seeking corrective surgery following a major life event
is likely to be a method of coping maladaptively with adjustment and loss; a characteristic presentation is the woman seeking breast augmentation after experiencing domestic violence and desertion by her partner. For this group of patients, psychiatric intervention can prove particularly helpful. Although psychiatric treatment has been advocated generally for dysmorphophobic patients in whom surgery is considered inappropriate, this is in practice rarely acceptable to the patient, and even when it is, successful interventions are the exception rather than the rule. Both serotonin-specific antidepressants and cognitive-behaviour therapy can be useful treatments for non-delusional body dysmorphic disorder.

**BORBORYGMI**

Harold Ellis

‘Borborygmi’ is the term applied to rumbling noises of varying quality and intensity produced by peristaltic movements of the bowel propelling mixed gaseous and liquid contents.

These sounds, although normally inaudible to the patient or to other persons, and detected only by auscultation by means of a stethoscope, may occasionally be annoyingly obtrusive. They may occur in perfectly normal people, especially when the alimentary canal is relatively empty, for instance when a meal is overdue, and they may occur as a result of nervous air-swallowing. They may be due to the powerful peristaltic waves of a bowel that is hypertrophied and dilated above a slowly developing obstruction of the large bowel; here there will usually be accompanying progressive constipation, colicky abdominal pains and distension. Some people are able to produce a loud sound by forcibly contracting the muscles of the anterior abdominal wall and splashing the fluid content of the stomach.

The carcinoid syndrome may feature loud borborygmi as well as flushing of the face, trunk and limbs, pulmonary stenosis, cramping abdominal pains and diarrhoea. In contrast, the absence of borborygmi, resulting in complete silence in the abdomen on auscultation for several minutes, is seen in adynamic ileus and peritonitis.

**BREAST, LUMPS**

Michael Baum

*(See also NIPPLE, ABNORMALITIES OF, p. 459.)*

**METHOD OF EXAMINATION**

The patient should sit stripped to the waist, so that a clear view of both breasts, the thorax, axillae and supraclavicular fossae may be obtained. The surgeon should sit with his or her eyes level with the nipples. Both breasts should first be looked at as a whole, to see whether they are symmetrical in size, contour and level, and whether the two nipples are in the same site and of the same circumference, prominence and inclination. One breast may always have been smaller, or one nipple inverted, but any recent change is highly significant. If no difference is at first noticed, the patient should be asked to raise both arms slowly above the head and bring them down again to the side, since differences previously invisible, particularly dimpling of the skin from attachment of a lump, may come into view as the breast glides over the chest wall. The patient should then lie on a couch, and the breasts be studied in detail for the evidence of local enlargement or shrinking, and for abnormalities such as redness of the skin, dilatation of the veins, a tumour or an ulcer. Next the breasts are felt, using first the flat of the hand, passing systematically over all parts; afterwards, the fingers are used for more detailed examination of any irregularity that may have been discovered or suspected. The axillae should also be palpated carefully for enlarged nodes, particular attention being paid to the inner wall, along the pectoralis minor and the apex. In cases of suspected cancer, the supraclavicular and infraclavicular fossae should also be examined for fullness or enlarged nodes, and the chest and liver should be investigated for signs of secondary growth. Examination from behind with the patient sitting may be used to check any abnormalities seen, felt or suspected in the lying position.

**Alternative posture for ‘difficult’ or pendulous breasts**

When dealing with a woman with large, obese or pendulous breasts, the conventional posture for examination is often unsatisfactory. An alternative posture is to arrange the woman in a semi-recumbent position, rotated obliquely with a pillow behind the scapula of the side under examination, and the shoulder fully abducted, with the hand tucked behind the head. This fixes the pectoralis major, and allows the breast disc to ‘float’ over a rigid base.

**CLASSIFICATION**

**Swellings of the whole breast**

- Bilateral
  - Pregnancy
  - Lactation
  - Diffuse nodularity (abnormalities of normal development and involution)
  - Hypertrophy
- In males, from drug or alcohol intake (oestrogenic agents and drugs for hypertension)
- Acute lactational mastitis

**Unilateral**
- Hypertrophy of the newborn
- Puberty
- Unilateral hypertrophy

**Discrete lumps in the breast**

**Benign**
- Fibroadenoma
- Simple cyst
- Galactocele
- Lipoma
- Plasma cell mastitis (periductal mastitis)
- Fat necrosis
- Tuberculous abscess
- Phyllodes tumour

**Malignant**
- Carcinoma
- (Rare: sarcoma, lymphoma)

**Multiple swellings, usually involving both breasts**

**Physiological changes with glandular hypertrophy**
**Multiple cysts**

**Swellings that are not of the breast**

- Retromammary abscess
  - From disease of rib
  - Chronic empyema
- Chondroma of chest wall
- Deformities of the ribs
- Mondor’s disease

**SWELLING IN PREGNANCY AND LACTATION**
Swelling in these cases is normal and only liable to cause confusion when the patient is unaware of her condition. Both breasts are enlarged equally and feel tense and nodular. The superficial veins are usually prominent and, on gentle squeezing, a few drops of milk can be discharged from the nipple. Montgomery’s tubercles will be evident.

**TRUE HYPERTROPHY**
True hypertrophy is rare. The enlargement is of two types: the more common in which multiple fibroadenomas cause a bilateral enlargement of varying consistency, and the less common consisting of a diffuse lipomatosis of both breasts, sometimes attaining prodigious proportions. The condition is usually bilateral, but it may be one-sided, in which case it is very disfiguring.

**UNILATERAL ENLARGEMENTS**
These are usually found in the undeveloped breast. In the newborn, one breast is often enlarged to the limits of its infantile size, and may discharge a little serous fluid from the nipple. The enlargement used to be attributed to manipulation by midwives, but it is more probably due to an endocrine imbalance consequent on the withdrawal of the maternal hormones in the fetal circulation, and it subsides rapidly. In girls at puberty, one breast may enlarge several months before the other and may distress a solicitous mother; unless there are obvious signs of an inflammatory change, no notice need be taken of unilateral enlargement of the breast in girls from 10 to 13. Uniform enlargement of one breast also occurs in men, usually after the age of 40, and nodular plaques may appear in both sexes at puberty as a result of endocrine disturbance.

On no account should the breast disc of an adolescent girl be biopsied, as this may cause failure of either a quadrant or the whole breast to develop, and this would be a legitimate reason for litigation.

**ACUTE MASTITIS**
Acute mastitis usually occurs during lactation, occasionally during pregnancy, and is most often due to infection with pyogenic organisms that have gained entrance through cracks in the nipple. At the beginning of the illness, there is shivering, followed by fever and a feeling of weight and pain in the breast; the pain soon becomes very acute. In the early stages, the swelling is limited to one part of the breast, which feels more resistant than normal; the skin is not reddened at first, nor are the lymphatic nodes enlarged. Pressure over the swelling may cause extrusion of a drop of pus from the nipple, and this is distinguished from milk by its viscidity and yellow colour. Later, fluctuation may become evident and, as the inflammation approaches the skin, this becomes red and oedematous, and ultimately an abscess may point and burst through it; at the same time, other foci of suppuration form, until the breast may be a bag of pus. The presence of fever and the intense tenderness of one portion of the breast are sufficient to distinguish acute mastitis from physiological engorgement.

It is not uncommon to find a small areolar abscess, which represents an infected gland of Montgomery.

**DUCT ECTASIA/PLASMA CELL MASTITIS (PERIDUCTAL MASTITIS)**
There is a common group of diseases that are generally poorly recognized and cluster together under this heading. Their aetiology is unknown. For example, it is even uncertain whether the inflammatory process comes first, followed by ectasia of the duct, or whether ectatic ducts are the primary phenomenon, with sloughing duct epithelium responsible for initiating the process of periductal mastitis. Assuming the latter sequence of events, the cycle of clinical features may
develop in the following way. The terminal lactiferous ducts dilate and often become hugely ectatic. As a consequence of this, the epithelial lining loosens and liquefies, causing plugs of cellular debris to fill up the ectatic ducts. The first clinical symptom of this condition is the extrusion of viscous multicoloured discharge from multiple duct orifices onto the nipple surface. The milk ducts then become permeable to the cellular and lipid contents normally contained within the lumina, and these then excite a chemical periductal inflammatory process, which is characterized by infiltration with plasma cells and foreign body giant cells. At this stage, a hard-indurated mass with overlying inflammation may appear at the areolar margin. Commonly, this condition resolves spontaneously within a week or two. Less often, the inflammatory mass becomes secondarily infected with anaerobic organisms liquefying to produce a periareolar abscess. This may point at the areolar margin, and spontaneously discharge. If the condition is not recognized and treated appropriately, a pathological communication between the ducts of the nipple and the skin develops, forming a so-called mamillary duct fistula. In the acute phase, antibiotics covering both aerobic and anaerobic organisms may abort the process. Over the years, a series of clinical or subclinical episodes of periductal mastitis produces fibrosis along the ducts, causing them to shrink and pull in the nipple, producing a typical slit-like indrawing at the centre. Ultimately, the condition burns itself out with age (Fig. B.10). This complex of conditions is most common in postmenopausal women but, if it occurs in premenopausal women, tends to be more florid and often bilateral, leading, to multiple abscesses and fistulae. Heavy tobacco consumption is now confirmed as a risk factor. The mammillary duct fistula should be treated by laying open the fistula track and excising the chronic inflammatory tissue. Surgically removing the whole of the subareolar system should treat recurrent episodes of periductal mastitis and troublesome nipple discharge.

**TUBERCULOUS ABSCESS**

Tuberculous abscess is rare in the Western world, but a certain number of cases of chronic mastitis and chronic abscess are really tuberculous, particularly in developing countries. The condition is also beginning to re-emerge in areas such as the East End of London, where there is a large influx of recent immigrants from Bangladesh. The disease is insidious, starting as a painless irregular swelling, the periphery of which is hard and the centre soft. Later, the skin becomes reddened, and an abscess forms, which may burst and leave a sinus. It differs from an acute abscess in that the duration is much longer, there is little or no pain or fever, and the pus, if examined, reveals no organisms on culture unless there has been secondary infection; a direct examination of stained films of the pus may show tubercle bacilli. The facts that the history is a long one, that the lump feels hard on palpation, and that the axillary nodes may be enlarged render this condition liable to be confounded with carcinoma.

**LOCALIZED FAT NECROSIS**

If this follows a blow to the breast, it may give rise to a tumour almost indistinguishable from cancer. It is hard, irregular in outline, and fixed to the skin. Points of distinction are the previous history of severe injury at the exact spot where the swelling lies, bruising and the impression given on palpation that the lump is on rather than of the breast, as well as the absence of hard nodes in the axilla. Sometimes a period of 2–3 weeks’ observation is justifiable, in which time a traumatic swelling should decrease in size, but if there is any real doubt about its nature, it should be excised and submitted to section. Contrary to popular myth, this is a rare condition of the breast. It is usually wise to ignore a history of trauma to the breast, which is often used by the woman as a rationalization for a lump, and fully investigate it with mammography and biopsy.
GALACTOCELE
A cyst containing milk, this is formed by dilatation of one of the larger ducts owing to obstruction. Galactoceles occur only during lactation and very rarely in the later months of pregnancy; they form oval, fluctuating swellings lying in the central zone of the breast just outside the areola and, on pressure, milk can sometimes be squeezed out of the nipple. Ultrasound scanning and aspiration both confirm the diagnosis and cure the condition.

SOLITARY CYSTS
Simple cysts are common in the 30–60 years age group. They present as well-defined, mobile lumps with a texture like that of an inflatable rubber ring. However, some are so tense as to be confused with a solid lump. Diagnosis is easy with a characteristic ultrasound image of a black, well-defined ovoid with a strong posterior acoustic enhancement. The diagnosis is confirmed and the condition treated by cyst aspiration. The cyst fluid should not be sent for cytological examination.

BENIGN TUMOURS
A fibroadenoma is the only common innocent tumour of the breast. It is an encapsulated lump, generally single but sometimes multiple, and varying in size. It is more common (and often multiple) in African-Caribbean women. It is firm, with the consistency of hard rubber, rounded or with irregular rounded projections, and clearly outlined. Most characteristic is the ease with which it can be moved under the skin and in the substance of the breast, to neither of which does it appear to have any attachment, hence the term 'breast mouse' that is applied to this lesion. These tumours generally occur between the ages of 18 and 30, and are quite painless. Although a carcinoma of the breast is rare in this age group, it is wise to complete the 'triple assessment' with an ultrasound scan and fine-needle aspiration cytology. If the diagnosis is confirmed by imaging and pathology, it is safe to reassure the young woman and discharge her from the clinic.

A lipoma may occur in the breast as elsewhere, and has the same clinical features as a lipoma in any other site. However, always beware the ‘pseudolipoma’ that may be the earliest sign of a small invasive duct cancer, which, by infiltrating Cooper’s ligaments, may extrude fatty lobules, forming a mushroom-like umbrella over the primary focus.

MALIGNANT TUMOURS
Malignant tumours of the breast are nearly always primary. Sarcoma is very rare, but carcinoma is common and the most important tumour that affects the breast. It is essentially a disease of the female breast, only about 1 per cent of the cases occurring in males. It is common in both married and unmarried women, and may occur at any age after puberty, although the majority are in women aged between 35 and 60. In advanced cases, the disease is obvious (Fig. B.11); the tumour is large and hard, attached to and ultimately, if not removed, fungating through the skin and becoming fixed to the chest wall; the axillary nodes are enlarged and hard. Such cases are beyond any but palliative treatment, and the importance of diagnosis lies in recognition of the early case, where the only sign is a small lump that the patient has probably discovered accidentally. Usually, there is no pain, and the patient looks and feels perfectly well. The lump may lie in any part of the breast, but typically is intermediate between the nipple and the periphery, and is more commonly in the upper and outer quadrant than in the other three. It can usually be felt with the flat of the hand. Such a lump may be stony hard, but any consistency may be met with. Its outline is usually not sharply defined. In the early stage, it is freely movable over the pectoral muscles and under the skin, but it is not so movable in the breast substance as is a fibroadenoma. Very soon, bands of fibrous tissue that connect the breast with the skin become involved, and by their contraction prevent free movement of the skin over the swelling; this causes first dimpling when the tumour is displaced, and later puckering visible all the time. If the tumour is anywhere near the centre of the breast, the nipple becomes retracted (Fig. B.12); a nipple may have been always inverted, but if one previously well formed becomes retracted, the sign is of serious import. Fixation to the deep fascia, which
usually comes later, can be demonstrated by making the patient press her hands onto her iliac crests to fix the pectoralis major, when the involved breast will be found to move less on the muscle than the normal one. Many cancerous tumours, even when extensive infiltration has occurred, cause shrinkage, so that the affected breast may appear smaller than the healthy one, and in the atrophic form it may almost disappear. In the ordinary form, it will be rare to find any discharge from the nipple. After a while, the axillary nodes become enlarged and hard. Too much attention should not be given to the absence of palpable nodes; in a fat patient, they may be enlarged but impalpable, and in any case it is hoped to recognize cancer before the nodes are involved.

**Duct carcinoma-in-situ**
The earliest premalignant condition affecting the breast ducts is referred to as duct carcinoma-in-situ (DCIS). Rarely the condition starts within the lobules, where it is referred to as lobular carcinoma-in-situ. In most cases, the condition is impalpable, and may only be discovered at a chance biopsy of a coincidental benign lump. More commonly these days, DCIS may be discovered as a result of a breast screening programme, where the condition shows itself as a cluster of microcalcifications on mammography, accounting for 20 per cent of screen detected ‘cancers’. Rarely, a large mass of DCIS of the comedo variety may present as a clinical mass or, if the in situ disease is close to the nipple, may present as a bloody nipple discharge or Paget’s disease (see NIPPLE, ABNORMALITIES OF, p. 459).

**Phyllodes tumour**
A phyllodes tumour is a rare clinical and pathological entity that presents with all the features of a giant fibroadenoma. In the past, this condition was referred to as a cystosarcoma phyllodes. However, the majority of these lesions are completely benign. The term phyllodes means ‘leaf-like’. This refers to the slit-like clefts arranged in a ‘botanical’ manner when viewed on a cut section. Rarely, the stromal elements of these tumours become hyperplastic and atypical, adopting some of the features of a sarcoma. This tumour is then referred to as a malignant phyllodes tumour. They have a tendency to recur locally if not widely excised at the first attempt and, with each recurrence, their malignant potential is more pronounced.

**ABNORMALITIES OF NORMAL DEVELOPMENT AND INVOLUTION (EXTREMES OF PHYSIOLOGICAL VARIABILITY)**
When a woman presents at the clinic complaining of a lump in her breast, the first step on clinical examination is to distinguish between a discrete lump and an area of lumpiness or nodularity. Although these lumpy areas in the breast of a young woman are extremely common (perhaps affecting 30 per cent of the female population), there is enormous confusion among the medical profession as to the correct terminology. In the past, these lumpy areas have been referred to by a series of terms, such as fibrocystic disease, fibroadenosis, mammary dysplasia, cystic hyperplasia, Schimmelbusch’s disease, chronic cystic mastitis, cystic mastopathy, König’s disease and masoplasia.

Whatever name is given to these lumpy breasts, there are no consistent pathological features that explain the varying textures palpable in different quadrants of the breast. For that reason, a group of clinicians in the University Hospital of Wales, headed by Professor Hughes, has come up with a rational description of these conditions, which can be grouped together under the catchy acronym ANDI, standing for abnormalities of normal development and involution. These abnormalities vary in extreme from normal physiological processes, which may be considered as benign disorders of little significance, to the other extreme, where the pathology can produce particular...
problems of discomfort or anxiety to the young woman. For example, during the developmental phase of the breast architecture, duct lobular overgrowth can lead to a fibroadenoma, which may cease growing at 2 cm or continue to grow to the extreme of a giant fibroadenoma, reaching sizes of 4 or 5 cm. Normal cyclical changes can produce premenstrual swelling and epithelial hyperplasia. Physiological abnormalities in sensitivity of the duct epithelium to the cyclical hormone changes can lead to cyclical mastalgie, nodularity and intraductal papilloma. Taken to its extreme, the intraductal epithelial hyperplasia can progress to atypia, which is known as a risk factor predicting the development of breast cancer. Finally, normal lobular involution may progress to the formation of cysts, sclerosing adenosis and duct ectasia. If a lumpy area of breast tissue is biopsied, almost all these features can be seen under the microscope to one extent or another.

Accepting that these conditions are aberrations of normal physiological development or involution, it can be accepted that, in most cases, the woman with a lumpy breast or a painful lumpy breast can be reassured. However, if the condition extends beyond the menopause into the cancer age group, clinical diagnosis can be extremely difficult, and it is in this area that ultrasound scanning and X-ray mammography are of great value. In addition, it should be remembered that invasive lobular cancer, which accounts for about 5 per cent of all malignant disease of the breast, may present as diffuse nodularity and atypical mammographic appearances. When in doubt, a core-cut biopsy is mandatory.

MULTIPLE CYSTIC DISEASE
Multiple cystic disease of the breast is usually regarded as a variety of ANDI. One breast, or sometimes both, becomes filled with cysts, some microscopic, others as large as walnuts, with all intermediate sizes, so that the organ has a bossy appearance. The diagnosis is usually simple, but it can be confirmed by aspiration of the cysts, a simple outpatient procedure that is also curative, although several aspirations may sometimes be necessary.

DIAGNOSIS OF A SINGLE LUMP IN THE BREAST
The diagnosis of a single lump in the breast, for which cancer must be taken into consideration, may cause considerable difficulty. A lump definite enough to be felt with the flat of the hand and hard enough to resemble cancer is a fibroadenoma, a tense cyst or a carcinoma. A fibroadenoma is usually found in women aged under 30, is less hard than a carcinoma and is of rounded outline, but its contour may be obscured by surrounding fibroadenosis. A cyst is usually round and elastic, but if it is deep its outline is obscured, and if it is tense it may feel hard. A carcinoma is undoubtedly solid and has an ill-defined outline; where these characters are present or where there is the slightest suggestion of skin dimpling, local flattening of the breast or alteration in the nipple, cancer must be diagnosed.

The diagnosis of cancer at this early stage is intensely important, for only then is the prospect of cure high. If there is the possibility that the lump is a cyst, this can easily be confirmed by ultrasound scanning; aspiration is then attempted under local anaesthetic. If clear fluid is obtained and the lump disappears, we can be certain that the diagnosis is one of a simple cyst. If no fluid, or only a few drops of blood, is obtained, smears should be made for cytological examination, a core-cut biopsy taken, or arrangements made for urgent excision and microscopic examination of the specimen. Local resection of a doubtful lump is imperative.

SWELLINGS PUSHING THE BREAST FORWARDS
These are often mistaken by the patient for breast tumours. A retromammary abscess is most commonly tuberculous, arising in an underlying rib or in a mediastinal abscess that has tracked along a branch of the internal thoracic artery. Sometimes an empyema points beneath the breast, usually in the fifth or sixth intercostal space in the mid-clavicular line. A chondroma is a hard nodular swelling springing from one of the ribs, and tilting the breast or pushing it aside. More common is a swelling of one or more of the costal cartilages, especially the second or third, which may be bilateral and tender. This condition, costochondritis or Tietze’s syndrome, is entirely benign and requires no treatment.

Deformities of the ribs may also cause confusion; the most common is a prominence of the costochondral junction of the third rib, which may be forked and join two cartilages. The condition is often bilateral, and may be associated with other abnormalities of the ribs or vertebrae.

ROLE OF X-RAY MAMMOGRAPHY AND ULTRASOUND SCANNING
It used to be widely accepted that routine X-ray mammography for women over the age of 50 who are otherwise asymptomatic might be of value in preventing premature death from breast cancer by the detection of subclinical cancers (Fig. B.13). This subject is now hotly debated and the pros and cons of screening are beyond the remit of this book. That aside,
no patient with breast cancer should be managed without mammography, as this will describe the extent of the disease within the ipsilateral breast and exclude the presence of synchronous contralateral cancers. Ultrasound scanning may help distinguish a solid from a cystic lump and may help define a discrete lump within an area of diffuse nodularity (see the section on ANDI, above).

MONDOR’S DISEASE

Although strictly speaking not a lump in the breast, it is difficult to know how to classify this condition. If a woman presents with characteristic guttering over the surface of the breast, this is a pathognomonic sign of Mondor’s disease, which is due to spontaneous thrombo-phlebitis of a superficial vein coursing over the thorax and breast. It has an indurated feel and can be mistaken for the dimpling of an underlying cancer, except for its linearity. The author has reported a case where Mondor’s disease was the only presenting sign of a cancer, which was detected on mammography. Other such cases have been reported, but it is unsure whether this may be coincidental or causal.

PRONOUNCED CYCLICAL MASTALGIA

Most young women notice some soreness and discomfort in the outer quadrants of their breasts during the week preceding a period, and this is a normal consequence of cyclical changes in their hormonal environment. In about 1 in 10 women,
this condition is sufficiently pronounced to cause anxiety, distress and insomnia. Characteristically, the pain is felt in the upper outer quadrants of both breasts, reaching a crescendo 2 or 3 days before the menses. Immediately after this, 2 weeks of comfort are experienced, and then the pain starts building up during the luteal phase of the cycle. The cyclical mastalgia may or may not be associated with lumpiness, and the two conditions should be considered separately. In the majority of such cases, simple reassurance with advice on mild analgesia is all that is needed. However, in severe cases, it is worth asking the woman to keep a diary, and if the pattern is clearly cyclical, a 6-month course of prolactin inhibitors (bromocriptine or danazol) can be prescribed. In addition, there is some circumstantial evidence that oil of evening primrose and the withdrawal of caffeine may benefit the condition, although it is notorious for this self-limiting disease to respond remarkably well to suggestion or placebo. By common consent, cyclical mastalgia has nothing to do with water retention, and therefore diuretics are not indicated. The underlying pathology is thought to relate to a hypersensitivity of the duct epithelium to biologically active prolactin.

Mammography is only indicated in women over the age of 35, or if pain is non-cyclical and localized to one area of the breast.

BRUISES

Mark Kiniron

Subcutaneous bleeding may present as bruises (ecchymoses), which will vary in colour from dusky red to green, yellow, purple or black depending on the duration of the bruise and the various haemoglobin breakdown products. Superficial crops of small capillary haemorrhages are called petechiae, and are best seen in the skin, mucous membranes and retina, although they also occur on the internal organs. Ecchymoses result from the confluence of petechiae or from haemorrhages from vessels larger than capillaries. Multiple bruises characterize bleeding disorders, whereas petechiae suggest an abnormality of either the platelets or the vessel wall. Multiple skin petechiae and ecchymoses are collectively referred to as purpura.

Bruising frequently occurs after trauma that may be minimal (or unobserved) in elderly persons, and isolated ecchymoses are normal in women and young children. There are several acquired vascular defects that promote skin bruising. Senile purpura affects older people, usually on sun-exposed areas such as the hands and forearms, and takes several weeks to resolve, leaving brownish discoloration (haemosiderin deposit). Corticosteroid use causes easy bruising and characteristic purple striae, particularly on the abdomen. Other vascular causes are vasculitis, connective tissue diseases and scurvy (a highly characteristic perifollicular haemorrhage, although gum bruising is also seen). Painful bruising syndrome occurs in women, and is associated with a tingling sensation followed by bruising over the trunk and limbs, with spontaneous resolution.

BULLAE AND VESICLES (BLISTERS)

Barry Monk

A blister is a circumscribed elevation of the skin containing free fluid. Blisters less than 5 mm in diameter are termed vesicles, while larger blisters are termed bullae. However, blisters of differing sizes may co-exist in some disorders, so the two presentations are best considered together. Blisters can be a feature of a number of important skin disorders (Box B.7).

TRAUMATIC DISORDERS

Normal skin is a cohesive, multilayered tissue that is remarkably resistant to friction, and only neonatal skin blisters easily. In adults, localized blistering of the skin occurs following thermal or chemical burns and sometimes frostbite. Friction due to ill-fitting footwear may cause blistering of the feet, while palmar blisters follow unaccustomed manual toil. Spontaneous blistering always has a pathological cause (Fig. B.14).

GENETIC DISORDERS

Rather rarely, a child is born with an inherited defect in the cohesion between the layers of the skin; electron
microscopy and genetic studies have now identified several forms of epidermolysis bullosa. The severest forms may be associated with mucosal (including laryngeal) involvement and severe blistering from birth, whereas other forms are associated with a rather milder degree of blistering, only occurring with friction. Another uncommon but important genetic cause of blistering in infants is incontinentia pigmenti, in which the linear pattern of blistering may lead the unwary to make the incorrect diagnosis of herpes zoster. This is an X-linked disorder, is generally fatal in the affected male, and is associated with a number of ocular and neurological disorders in affected females.

INFECTION DISORDERS
A number of infections can cause blistering eruptions in infants and young children. Bullous impetigo (Fig. B.15) is caused predominantly by superficial infection with Staphylococcus. Sometimes the predominant lesion is the honey-coloured oozing crust, but quite marked blistering may occur. The roof of the blisters is fragile and easily ruptured, leading to an appearance that has been likened to a cigarette burn, and that may lead to false accusation of child abuse or neglect. Microbiology swabs will demonstrate the causative organism, and there is a rapid response to topical antibiotics. Some strains of Staphylococcus produce an exotoxin that causes a dramatic superficial exfoliation of the skin resembling that caused by a burn (staphylococcal scalded skin syndrome); this must be distinguished from toxic epidermal necrolysis, a condition that may arise at any age, usually as a result of an idiosyncratic reaction to a drug, in which extensive areas of skin apparently dissolve leaving raw weeping areas; the mortality is high.

Another cause of blistering in young children that may cause diagnostic confusion is scabies; this is an infection of the skin with the mite Sarcoptes scabiei, which is passed from person to person by direct physical contact. The classic lesion of scabies is the burrow; this is an irregular serpiginous track that is most commonly found around the wrists or finger webs, and from which the live mite may be extracted on the end of a needle and demonstrated under the microscope. In babies, blistering on the soles of the feet may be found and should prompt an examination of other family members (especially the mother).

The sudden onset of a widespread vesicular rash, associated with mild fever and constitutional upset, may suggest chickenpox (varicella; Fig. B.16). Blisters appear in crops, so will be at different stages of evolution, and mucosal blistering may be evident in the mouth. In patients with any form of immunodeficiency or who are receiving immunosuppressive drugs, chickenpox may present with a violent haemorrhagic blistered eruption; pulmonary involvement may occur, and there is a significant mortality.

The cropped eruption of chickenpox is one of the features that was used to distinguish it from smallpox (variola), a condition now thankfully eradicated, in which the lesions erupt together and are thus much more monomorphic. A generalized vesicular or pustular eruption is a rather uncommon sequel to smallpox vaccination.

Perhaps the most common of all infections to cause blistering is Herpes simplex (Fig. B.17). The typical...
lesion of herpetic infection is a cluster of small vesicles, arising on an erythematous background, which crust and heal over a period of 7–10 days, only to recur repeatedly on the same site. When herpetic vesicles occur on mucosal surfaces, the appearance may be somewhat modified; vesicles become macerated and the roofs rub off, leaving painful round erosions or shallow ulcers. The sites most commonly affected are the lips, the face, the buttocks and the genitalia. Although genital herpes simplex (type II herpes) typically occurs on the genitalia, type I infections may also arise at this site. The first episode of herpes simplex is usually more severe, and is associated with more pain and constitutional upset than subsequent episodes.

The history of recurrent blistered lesions at the same site is characteristic of herpes, but it is also the hallmark of another blistering disorder, fixed drug eruption. In this dramatic and intriguing condition, a blister on an erythematous base recurs in exactly the same location (commonly including the orogenital mucosae) each time affected individuals are exposed to certain medicaments (e.g. phenolphthalein laxatives, codeine, tetracyclines or sulphonamides). The blister resolves, leaving a round patch of post-inflammatory pigmentation, which will then be the site of the next episode. It is remarkable how rarely patients associate the recurrent painful blistering with taking the offending medication.

Herpes simplex, although usually causing a localized blistering, may be responsible for a widespread eruption in patients with atopic eczema, who appear to be incapable of mounting an appropriate immune response to the virus, eczema herpeticum (Kaposi’s varicelliform eruption). Typically, the subject has quite active eczema, and presents with a sudden onset of a diffuse or generalized eruption composed of umbilicated vesicles or crusted erosions. The lesions are often painful, and there may be a severe fever and constitutional disturbance. The condition may be fatal, especially if unrecognized and appropriate treatment not instituted.

Herpes zoster (shingles) causes a band of vesicles along a dermatome. It is caused by a secondary activation of varicella virus lying dormant in nervous tissue following primary (chickenpox) infection, sometimes as long as 60 years previously. The condition is uncommon in childhood (except, curiously, in black individuals) and more common with increasing age. It also occurs in those immunosuppressed by drugs or HIV disease, systemic illness or malignancy, where it may be accompanied by toxicity and disseminated chickenpox lesions. Pain usually precedes obvious skin changes by up to 48 hours. The first sign in the skin is erythema on which grouped clear vesicles appear, which later become umbilicated and haemorrhagic. Smears from the base
The vesicles show multinucleated giant cells. The limitation to one side of the body, the distribution in one or more dermatomes and the pain usually suffice to distinguish herpes zoster from erythema multiforme and dermatitis herpetiformis (Fig. B.18). When the trunk is affected, the preceding pain may be mistaken for pleurisy or some intra-abdominal condition. The pain may also simulate myocardial infarction, a slipped disc, orchitis or venous thrombosis depending on the site affected. When the second division of the trigeminal nerve is affected, involvement of the tip of the nose with vesicles indicates infection of the nasociliary branch of the ophthalmic nerve and hence is a sign that keratitis may occur.

Sometimes, the bites and stings of insects cause bullous reactions and present as discrete large blisters on the lower legs, especially in children (Fig. B.19). The sting of jellyfish produces a bullous reaction. In dermatitis artefacta, bullae may be artificially induced by patients themselves (e.g. with caustic chemicals). Blister can be a feature of acute severe eczema of any cause (Fig. B.20). This is seen on the palms, sides of fingers and soles in 

\[ \text{pompholyx} \]

and in acute contact dermatitis. Some plants, especially of the Umbelliferae family, produce a toxin that is activated by ultraviolet light. Typically, it presents in gardeners who have been clearing a patch of overgrown land on a sunny day, and wearing short sleeves or short trousers.

**INFLAMMATORY DISORDERS**

Another condition characterized by blisters provoked by sun exposure is porphyria cutanea tarda; small vesicles occur on the backs of the hands and are associated with an acquired fragility of the skin. The urine will fluoresce when examined under ultraviolet light. Blistering on the legs may occur in patients with renal failure on haemodialysis; its cause is unknown.

**Erythema multiforme** (Fig. B.21) is an acute eruption that may be triggered by a large number of provoking factors, including viral infections, drugs and radiotherapy. Recurrent episodes occur if the provoking cause recurs (e.g. herpes simplex infection). The characteristic ‘target’ lesions on the dorsa of the hands and feet, over the knees and elbows, and sometimes
more widely, owe their pattern to two erythematous rings of different shade surrounding a central blister. In severe cases, blistering may be extensive and associated with fever, prostration and occasionally pneumonitis. Involvement of the mucous membranes may be a prominent feature (Stevens–Johnson syndrome; Fig. B.22).

**IMMUNOBULLOUS DISORDERS**

Three uncommon conditions – pemphigus, pemphigoid and dermatitis herpetiformis – share the features of being chronic blistering eruptions, in which characteristic patterns of deposition of antibodies in the skin can be demonstrated.

**Pemphigus** comprises a group of disorders, possibly arising in those with a genetic predisposition, in which there is formation of blisters within the epidermis. Both the skin and mucous membranes are affected. **Pemphigus vulgaris** begins in the third and fourth decades of life, with equal gender-related incidence. The mucosa of the mouth is often affected before the skin. Fragile intra-epidermal cutaneous blisters occur at any site and quickly become erosions. These are often extensive and slow to heal, and become secondarily infected. The uncomfortable oral lesions are the most constant, persisting even when blistering elsewhere is controlled, and making eating difficult. The blistering is thought to be due to a circulating autoantibody directed against the intercellular substance. This is detectable in the blood by an indirect immunofluorescence technique on a suitable substrate tissue (e.g. primate oesophagus), and the titre of this antibody is related to the severity and progress of blistering. Fluorescence may also be demonstrated in unfixed skin biopsy specimens. Affected patients also have a higher incidence of organ-specific autoimmune diseases and thymoma. A milder form of pemphigus with positive immunofluorescence may arise in patients under treatment with penicillamine. **Chronic familial benign pemphigus** (Hailey–Hailey disease) is unrelated but has histological similarities. Affected family members show a vulnerability of flexural skin to friction, producing characteristic fissured erosions on the sides of neck, axillae, perineum and oral and vulval lips. Circulating autoantibodies have not been detected.

**Bullous pemphigoid** (Fig. B.23) is usually a disease of the elderly. Bullae of 2–5 cm diameter arise on an erythematous, urticated background on the limbs and trunk. The blisters are thick-roofed and can last for many days; initially, their constituent fluid is clear, but it soon becomes cloudy or haemorrhagic. Once the multiple large blisters occur, the diagnosis can be easily made. However, widespread pruritus, erythema and urticaria may precede the blistering by several weeks or
months, and at this stage diagnosis is difficult. There is also a circulating autoantibody, but in this case directed at the dermo-epidermal junction. The mucosae can be affected with similar tense bullae, but this is a much less constant feature than in pemphigus. 

*Cicatricial pemphigoid* (benign mucous membrane pemphigoid; Fig. B.24) is a variant in which the blister forms just below the dermo-epidermal junction, so that healing takes place with scarring. Blisters appear on mucosae rather than skin, particularly the eyes, frequently causing blindness.

In *dermatitis herpetiformis*, the blisters are subepidermal and tend to be short-lived, as, being intensely pruritic, they are quickly excoriated so the diagnosis may often be missed for months or years. The condition chiefly affects the extensor surfaces of the shoulders, buttocks, knees, forehead and scalp, sparing the mucosae. There appears to be a related gastrointestinal gluten hypersensitivity, and direct immunofluorescence of frozen skin reveals diagnostic deposits of IgA in the dermis. The condition responds dramatically to dapsone, its mode of action being uncertain.

**Figure B.24** Cicatricial pemphigoid.
CATARACT
Reginald Daniel

A cataract is an opacity within the lens of the eye; the opacity may be located in the centre of the lens (nuclear), in the cortical region or in the posterior subcapsular area. It has little clinical significance unless interference with vision results. Cataracts are frequently bilateral, but the severity and rate of progression in each eye usually varies. Most cataracts are associated with ageing, but they can result from a wide variety of causes. Cataracts may have a congenital or acquired aetiology and sometimes occur secondary to other eye diseases such as iritis, glaucoma and retinitis pigmentosa. Cataracts can result from trauma and are often associated with some systemic disorders (e.g. diabetes mellitus) and certain drugs used topically or systemically (e.g. corticosteroids) (Box C.1).

Most cataracts are not visible to the casual observer until they are advanced and causing profound visual loss (Fig. C.1). In the early stages, cataracts are best diagnosed by examining the red reflex of the fundus with an ophthalmoscope through a well-dilated pupil, or by slit-lamp examination. The most common symptom caused by cataract is decreased visual acuity. Other symptoms include increasing myopia, monocular diplopia, and glare in which vision is much worse in conditions of bright illumination.

The rate of progression of cataract is very variable; when the visual loss is interfering significantly with the patient’s daily living, surgery is indicated. The method of cataract extraction surgery has advanced considerably over the years. The modern technique is phacoemulsification, in which the lens with the cataract is emulsified using an ultrasonic handpiece through a small incision, and the lens particles are aspirated from the eye. This procedure is usually performed under topical or local anaesthesia, but general anaesthesia is required for children and very apprehensive or uncooperative patients.

Lens extraction with the insertion into the lens capsular sac of an intraocular lens implant manufactured from polymethylmethacrylate, silicone or acrylic material restores vision very successfully. On rare occasions, it may not be possible to implant an artificial lens, and in these cases either contact lenses or cataract spectacles are necessary to attain a satisfactory result. Intraocular lens implants have become extremely refined and usually achieve excellent postoperative vision with a very low complication rate. Implants have considerable optical advantages over cataract glasses, which are associated with disturbing problems of image magnification, lens aberrations and limited visual field. Contact lenses overcome these optical problems, but many elderly or handicapped patients experience management and handling problems.

CHEST, DEFORMITY OF

Box C.1 Causes of cataract

- Age-related (senile) cataract
- Cataract associated with ocular disease
  - Congenital disorders
  - Acquired disorders
    - Uveitis
    - Glaucoma
    - Neoplasia
    - Topical drug therapy – steroids and miotics
    - Trauma – mechanical, chemical or radiation
- Cataract associated with systemic disease
  - Maternal infection
    - Rubella
    - Cytomegalovirus
  - Hereditary disorders, e.g. Down’s syndrome, dystrophia myotonica, Alport’s syndrome, Fabry’s disease, Lowe’s syndrome
  - Metabolic disorders, e.g. diabetes mellitus, galactosaemia, mannosidosis, hypoparathyroidism, hypothyroidism
  - Systemic drugs, e.g. corticosteroids, antimiotics, phenothiazines and anticholinesterases
  - Dermatological disorders, e.g. atopic dermatitis and ichthyosis
no significant effect on respiration may cause concern to the patient and are relatively common. These are described, together with the infrequent but more serious conditions that adversely affect cardiorespiratory function.

The normal configuration of the chest is influenced by the age, sex and physical build of the individual. It is determined by the condition of the spine, ribs and sternum, the overlying muscles and soft tissues, and the underlying lung and pleura. In infants, the chest wall is almost circular in cross-section, the ribs lie horizontally, and the anteroposterior and transverse thoracic diameters are similar. With growth, the chest becomes flattened anteroposteriorly and wider transversely, and the ribs adopt an oblique, downward-sloping position.

The shape of the adult chest is dependent upon body build. In stocky mesomorphic individuals, the chest wall tends to be circular, with relatively deep posteroanterior and wide transverse diameters. The heart may lie horizontally. The vertical height from the sternal notch to the diaphragm is proportionally reduced. In contrast, the chest in ectomorphic individuals is long in proportion to the overall width and shallow anteroposteriorly, and the heart adopts a more vertical position (Fig. C.2). Variation in normal body build is not associated with a predisposition to respiratory disease, although morbid obesity may cause respiratory failure, either directly by influencing chest wall function or indirectly by provoking obstructive sleep apnoea.

CONGENITAL DEFECTS

Rib abnormality
Bifid ribs are common, particularly in the upper six ribs, and may cause confusion if the abnormality is not appreciated on the chest radiograph.

Cervical ribs, usually arising from the seventh cervical vertebra, occur in 0.5 per cent of the population; in 80 per cent, they are bilateral, and they vary greatly in size and shape. They are seldom a cause of symptoms, and are often a chance discovery either on a routine chest radiograph or at physical examination when a deep bony mass is discovered in the supraclavicular fossa. Symptoms due to compression are more prevalent in females, and more common on the left side. Neurological symptoms include pain and weakness in the arm, with paraesthesiae of the fingers and wasting of the intrinsic muscles of the hand. Vascular symptoms may mimic Raynaud’s phenomenon, and subclavian artery obstruction or thrombosis may cause distal gangrene.

Pectus carinatum or pigeon-chest deformity
The sternum is prominent, forming an anterior ridge, and the ribs are inclined forwards, causing a greatly increased anteroposterior diameter. The condition can be acquired (asthma), but if congenital is due to premature obliteration of the sternal sutures during growth, or to malattachment of the anterior portion of the diaphragm to the posterior portion of the rectus sheath rather than, as normally, to the xiphoid process, with consequent distorting mechanical effects.

Pectus excavatum or funnel-chest deformity
The costal cartilages are prominent and curve inwards, and the body of the sternum is depressed backwards towards the spine from the manubriosternal joint downwards, with maximum recession at the xiphoid. In severe cases, the lower sternum forms a deep concavity, and may almost touch the spine. The heart is displaced to the left side of the chest. Radiographic displacement of the heart to the left, and rotational changes of the electrocardiogram (ECG), may be wrongly interpreted as evidence of heart disease. The ECG may show persistence of the juvenile pattern, with T wave inversion in the right precordial leads, incomplete right bundle-branch block and right axis deviation.

Figure C.2 Radiograph of a normal chest. Patient of tall, thin build with a vertically disposed heart.
Because of cardiac rotation, there may be P wave inversion and a QR pattern in lead VI. Minor lung function abnormalities occur, with reduced total lung, maximum breathing and vital capacities. The condition does not predispose to cardiac or respiratory disease in later life. Surgery is rarely required because of symptoms, but it is occasionally sought for cosmetic reasons.

**Incomplete fusion of the sternum**
This is an unusual abnormality, apparent at birth, producing the appearance of a split sternum with indrawing of the soft tissue over the central fissure during inspiration and bulging on expiration. This paradoxical respiratory movement is much increased when coughing or in the presence of respiratory obstruction.

**Partial or complete absence of the pectoral muscle**
Such an absence is a rare congenital abnormality, usually unilateral and mostly involving the lower portion of pectoralis major. The condition produces no symptoms, but if it is of a severe degree, the rib cage is deformed and the anterior chest wall on the affected side is underdeveloped and shrunken because it is not subject to the lateral pull of the pectoral muscle. The chest radiograph may show abnormal transradiancy of the affected side, which may give rise to an erroneous impression of pulmonary disease.

**The straight back syndrome**
This is an absence of the normal physiological dorsal kyphosis of the spine associated with a reduced anterioposterior diameter of the chest. A mild insignificant restrictive defect of lung function may be present, but cardiac complications are more likely to occur. Examination may reveal a palpable left parasternal systolic impulse and exaggerated splitting of the second heart sound on auscultation, presumably caused by compression of the pulmonary outflow tract and great vessels between the spine and sternum. The ECG may show an RSR pattern in lead VI.

**ACQUIRED SKELETAL DEFORMITIES OF THE CHEST**

**Scoliosis**
(see also SPINE, DEFORMITY OF p. 627)
Scoliosis is a lateral curvature of the spinal column and may either be functional or structural. A functional or non-structural scoliosis is one in which the curve is correctable on sitting or lying. The commonest cause for this is a leg-length discrepancy leading to a spinal tilt that is relieved by sitting. A structural scoliosis is accompanied by rotation of the apical vertebral bodies into the concavity of the scoliosis. It is this combination of lateral displacement and rotation which gives rise to the deformity of a rib hump and anterior chest wall asymmetry (Figure C3).

The causes of a structural scoliosis are multifactorial, the commonest type being idiopathic scoliosis, which is a curve for which no discernable cause is able to be determined, but it is almost certainly genetic in origin. The other causes are listed in Box C.2.

*Figure C.3* This illustrates a patient with adolescent idiopathic scoliosis with a right rib hump which becomes more apparent with the Adam's forward bending test as shown.
Idiopathic scoliosis is considered as early onset if it occurs before the age of 7 years of age and late onset after the age of 7 years. The age presentation was previously divided into infantile – 0 to 3 years, juvenile – 3 to 10 years and adolescent - after 10 years of age to reflect the different periods of growth that occur in children. The reason for the change to early and late onset is to reflect the pulmonary growth, in that the majority of the alveola number and volume are developed by the age of 7 years. Thus, if a spinal deformity develops before the age of 7 years, it causes a significant decrease in the chest volume, significantly inhibiting pulmonary development leading to significant respiratory compromise known as thoracic insufficiency syndrome. If the scoliosis develops after the pulmonary tree has developed (Fig C.4), there is significantly less impact on respiratory function and it is only when curves are over the size of 100 degrees that any significant respiratory compromise occurs and it only then leads to mild loss of function for the patient.

**Box C.2 Causes of scoliosis**

<table>
<thead>
<tr>
<th>Congenital</th>
<th>Idiopathic</th>
<th>Neuromuscular</th>
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<tbody>
<tr>
<td>Failure of segmentation, formation or both, of the vertebral body leading to scoliosis</td>
<td>Early Onset</td>
<td>Neuropathic</td>
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<td></td>
<td>Late Onset</td>
<td>UMN, Cerebral Palsy</td>
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<td>Polio, Spinal Muscular Atrophy</td>
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<td>Neurofibromatosis</td>
<td>Myopathic</td>
<td>Fibre type disproposition</td>
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<td>Mesenchymal disorders</td>
<td>– muscular dystrophies, eg Duchenne’s</td>
<td>Congenital Hypotonia</td>
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<td>Marfan’s syndrome</td>
<td>– fibre type disproposition</td>
<td>Myotonic Dysrophies</td>
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<td>– congenital hypotonia</td>
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<td>– costochondral dysplasia</td>
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</tr>
<tr>
<td>Sepsis</td>
<td>– muscular dystrophy</td>
<td></td>
</tr>
<tr>
<td>Metabolic</td>
<td>– muscular dystrophy</td>
<td></td>
</tr>
<tr>
<td>Tumours</td>
<td>– muscular dystrophy</td>
<td></td>
</tr>
</tbody>
</table>

**Figure C.4** Whole-spine radiograph of a young man with idiopathic scoliosis.

**Ankylosing spondylitis**

This is an inflammatory arthritis affecting the axial skeleton of young men, the great majority of whom are HLA-B27 positive. The condition results in the fusion of the costotransverse and vertebral joints, with relative fixation of the rib cage in an inspiratory position. The sternomanubrial and sternoclavicular joints may also be affected. Clinical examination of the chest may reveal a dorsal kyphosis and diminished chest expansion, with a corresponding increase in abdominal expansion. Changes in the relative proportion of diaphragmatic and chest wall movement alter the normal relationship of ventilation to perfusion in the upper lobes. Most of the ventilatory movement is performed by the diaphragm and abdomen. With advanced disease, signs of apical fibrosis, consolidation or cavitation may be present. The posteroanterior chest radiograph may demonstrate the characteristic bilateral apical cavitating lesions, and the lateral radiograph the classical ‘bamboo’ calcification of the anterior spinal ligaments. The clinical picture is completed if there is an aortic diastolic murmur and cardiac failure due to aortic valve disease.
Rickets (osteomalacia)
Rickets, either nutritional or due to renal disease or inherited enzyme defects (1-alpha-hydrolase), may cause important skeletal deformity, with disorganization of the bone growth plate and faulty replacement of calcified cartilage. In active childhood rickets, the costochondral junctions enlarge, producing the so-called 'rickety rosary'. The soft, plastic ribs are readily moulded. The upper ribs tend to be indrawn by inspiratory forces, while the lower chest is supported at the costal margins by the underlying abdominal viscera. A depression is produced above the costal margins, with a flaring outwards of the costal margin itself. This deformity may remain and is known as Harrison's sulcus.

Pigeon chest
This condition may be congenital (rare), due to rickets or most commonly accompanies chronic bronchopulmonary disease in childhood. This is most frequently seen as a skeletal manifestation of severe uncontrolled asthma. It is, to a considerable extent, reversible if the asthma is treated adequately and sufficiently early. It should be a rarity. Pigeon-chest deformity may also occur in bronchiectasis.

Barrel chest
Barrel chest is the name given to the deep chest in which the anteroposterior diameter is increased, often to equal the transverse diameter, the subcostal angle is abnormally wide, and the horizontal position of the ribs is accentuated. This appearance is most constantly related to overinflation of the lungs, and may be observed in acute attacks of asthma and in chronic obstructive pulmonary disease, usually associated with irreversible structural emphysema, especially when associated with alpha-1-protease deficiency. Radiologically, the lungs are hyperinflated and hypertransradiant, the heart is narrow and lies vertically, and the diaphragm adopts a low position.

General expansion of one side of the chest
General expansion of one side of the chest is unusual. It may occur in children and young adults but is unlikely in older subjects with more rigid chest walls. A clinically detectable expansion of one hemithorax may accompany large rapidly accumulating pleural effusions, large tension pneumothoraces, especially in patients undergoing assisted ventilation, or unilateral obstructive emphysema associated with partial bronchial obstruction, causing a check valve effect that allows air to enter the lung segment but prevents its escape, and with an overexpansion of giant air-containing cysts in the lung.

General contraction of one side of the chest
General contraction of one side of the chest is due to a loss of volume of the underlying lung. This may follow lung collapse, lung fibrosis, surgical resection of the lung tissue or pleural thickening preventing inflation of the underlying lung. Pleural thickening may result from chronic empyema (Fig. C.5) (bacterial and tuberculous), or malignant...
infiltration from adenocarcinoma or mesothelioma. Unilateral pulmonary fibrosis may result from lung inflammation due to pulmonary tuberculosis, chronic lung abscess or organizing pneumonia. The whole hemithorax may appear shrunken, with abnormalities most apparent in the upper anterior chest, showing obvious flattening and reduced respiratory excursion (remember the adage ‘flattening equals fibrosis’). The contracted fibrotic lung due to chronic pulmonary suppuration is frequently the site of bronchiectasis (Fig C.6).

When unilateral contraction of the chest with gross pleural thickening arises from any cause during childhood or adolescence, the resulting scoliosis may become extreme as growth proceeds.

**Table C.1 Causes of localized swelling of the chest wall**

<table>
<thead>
<tr>
<th>Infectious inflammations</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Tuberculosis of the chest wall</td>
<td>See text</td>
</tr>
<tr>
<td>Osteomyelitis</td>
<td>• In children: staphylococcal infection involving the ribs</td>
</tr>
<tr>
<td></td>
<td>• In adults: usually staphylococcal in drug addicts following a blow</td>
</tr>
<tr>
<td>Osteochondritis of costochondral junction</td>
<td>• Tietze’s syndrome (uncommon); see text</td>
</tr>
<tr>
<td></td>
<td>• Typhoid fever (very rare); see text</td>
</tr>
<tr>
<td>Syphilitic gumma</td>
<td>Starts in anterior mediastinum and presents at the front of an intercostal space. Now very rare</td>
</tr>
<tr>
<td>Actinomycosis</td>
<td>See text</td>
</tr>
<tr>
<td>Fungal infections</td>
<td>Blastomycosis, coccidiomycosis and cryptococcosis can cause osteolytic rib lesions</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Benign soft-tissue tumours</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Lipoma</td>
<td>Most common benign tumour of the chest wall. Computed tomography scan confirms the presence of fat</td>
</tr>
<tr>
<td>Haemangioma</td>
<td>Mass may pulsate. Bruit heard. Phleboliths in the tumour on chest radiograph diagnostic</td>
</tr>
<tr>
<td>Cystic hygroma</td>
<td>A rare hamartomatous lymphatic malformation producing a thin-walled cystic tumour lined with endothelium</td>
</tr>
<tr>
<td>Neurofibromatosis</td>
<td>Multiple fibromas present in one or more intercostal spaces. Café-au-lait patches, etc.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Malignant soft-tissue tumours</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Fibrosarcoma</td>
<td>Most common malignant soft-tissue primary tumour of the chest wall; can arise in soft tissues in older women many years after radiotherapy to the breast</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Benign bony tissue tumours</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Osteochondroma (exostosis)</td>
<td>Small hard tumour fixed to a rib or the sternum in children and young adults. May cause severe pain</td>
</tr>
<tr>
<td>Chondroma</td>
<td>Tends to become carcinomatous; see Chondrosarcoma</td>
</tr>
<tr>
<td>Diaphysiological aclasias</td>
<td>Generalized disorder of skeleton. The rib excrescences resemble chondromas</td>
</tr>
<tr>
<td>Osteoclastoma</td>
<td>Rarely affects ribs. About 10% can become malignant</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Malignant bony tissue tumours</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Metastases</td>
<td>Primary is usually in lung, breast, prostate, thyroid or kidney in adults, and is usually a neuroblastoma in children</td>
</tr>
<tr>
<td>Multiple and solitary myeloma</td>
<td>Monoclonal hypergammaglobulinaemia</td>
</tr>
<tr>
<td>Chondrosarcoma</td>
<td>Initial silent painless growth – may stop growing for a time only to become painful and invasive</td>
</tr>
<tr>
<td>Ewing’s sarcoma</td>
<td>Aggressive, most common in those under 30 years of age. Involves the ribs and rarely scapula or sternum</td>
</tr>
<tr>
<td>Osteogenic sarcoma</td>
<td>May occur in rib of patient with Paget’s disease or in a woman many years after radiation therapy to the breast</td>
</tr>
</tbody>
</table>
chest wall. Abscesses may arise from nodes in the region of the internal mammary artery, from which pus may present near the costochondral junction. Tuberculosis of the sternum can cause a cold abscess over the anterior chest wall. Rib tuberculosis is rare but may give rise to local pain, swelling and sinus formation. Radiologically, there is an initial small area of bone destruction that progresses to periosteal elevation and soft-tissue swelling. Multiple cold abscesses of the ribs may accompany spinal tuberculosis (Pott’s disease), with paravertebral abscess formation pointing over the back and lower ribs.

**Tietze’s syndrome (costochondritis)**

This condition of obscure aetiology results in pain, swelling and tenderness of one or more of the upper six costal cartilages. The second costal cartilage is most often affected. It may present at any age, but it is most common in young adults. The onset may be insidious or abrupt, the pain variously described as aching, gripping, sharp or dull, and the condition, which is self-limiting, can persist for weeks, months or rarely years. It has been suggested that asymmetrical rib growth, prolonged coughing or hyper-ventilation may be the cause, but it is often idiopathic.

**Osteomyelitis of the ribs, spine or sternum**

Osteomyelitis is a well-known, albeit rare, consequence of bacteraemia. Classically difficult to diagnose, patients present with chronic symptoms of pain, fever, thoracic cage swelling and radiographic evidence of erosion and bony sclerosis. *Staphylococcus aureus* is the most common agent. *Staphylococcus epidermidis* has emerged in recent years as a frequent cause from infected intravenous lines and prosthetic implant material. Gram-negative organisms are often responsible for haematogenous vertebral osteomyelitis, especially in patients with sickle-cell anaemia. Intravenous drug abusers are especially susceptible to a wide range of bacterial and fungal causes of osteomyelitis.

**Actinomycosis**

Thoracic actinomycosis may spread from the lungs to cause a diffuse indurated swelling of the chest wall; if the condition goes undiagnosed and untreated, it progresses to destructive rib lesions associated with periostitis and multiple sinuses discharging ‘sulphur granule pus’.

**Empyema necessitatis**

A localized swelling of the chest wall may be caused by a neglected empyema pointing externally to produce a soft, diffuse, fluctuant swelling that sometimes gives an impulse on respiration or coughing.

**Tumours**

Tumours may originate or metastasize to the chest wall (see Table C.1).

**Aortic aneurysms**

Aortic aneurysms involving the ascending part of the aorta may cause pulsating swellings over the upper anterior chest; they should be easily recognizable by the characteristic expansile pulsation, and by other signs and symptoms of an aneurysm. The most common situation for such a swelling is to the right of the sternum in the first, second and third intercostal spaces. In the presence of a grossly enlarged heart, the precordium may become prominent, a condition most often seen in children suffering from severe rheumatic or congenital heart disease.

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**CHEST, PAIN IN**

Alex West

Pain in the chest is one of the most common of all complaints. However slight it may be, it only too often conjures up in the mind of the sufferer a vision of serious organic disease of the lungs or, more often, of the heart. Accurate diagnosis depends, to a considerable extent, on physical examination and special investigations but most of all on a detailed and precise history of the site and radiation of the pain, of its character and duration, and of factors by which it is aggravated or relieved. A useful clinical classification of pain in the chest is into pain felt mainly in the centre of the chest or in its lateral aspects; each of these categories can be further subdivided into: (i) pain of sudden onset, often presenting as an emergency; and (ii) pain which, on presentation to the physician, has been present for several days or weeks. The common, and some of the less common, causes of chest pain are listed in this way in Table C.2. Pain in the precordium will be included in the discussion of central chest pain. The adjectives ‘central’ and ‘precordial’ are not of course synonymous – the precordium being that area of the anterior chest wall circumscribed by the surface markings of the heart – but the two sites are naturally associated in the diagnosis of pain that arises, or is thought to arise, from the heart. Before proceeding to the main discussion, a brief account will, for the sake of completeness, be given of several causes of chest pain that are immediately obvious on superficial examination.
PAIN DUE TO SUPERFICIAL LESIONS

Pain due to inflammation of the superficial tissues of the chest wall poses no great diagnostic problem. It is important to remember, however, that the inflammation may have spread from a deeper lesion such as an empyema. Herpes zoster, which involves thoracic nerve roots in at least 50 per cent of cases, is also obvious once the eruption has appeared, but pain and paraesthesia may be present for a few days before this and cause temporary diagnostic confusion. The vesicles are implanted on an erythematous base and may be discrete or confluent; they are strictly unilateral and occupy the area of one or more dermatomes. Fever and malaise occur in a few patients, and the thoracic nerve roots in at least 50 per cent of cases, is other scapular region. The pain is described as ‘tight’, ‘gripping’ or ‘like indigestion’, or the patient may deny actual pain and describe only a feeling of pressure or of tightness. The patient may place a clenched fist placed on the sternum to indicate both the site and, presumably, the character of the sensation (Levine's sign). A pain described as ‘stabbing’ is probably not angina, but patients do not always choose their words with care, and each statement must be carefully analysed to ascertain exactly the patient's meaning. Certainly, a pain that comes in sharp jabs, lasting a second only, is unlikely to be angina, and adjectives such as ‘shooting’ and ‘stabbing’ are suspect on this account. The typical duration of an attack is only a few minutes, and a pain lasting for a much longer or shorter time than this is unlikely to be angina; there are exceptions, however, which will be discussed. Perhaps the most important aspect is the relationship to exertion. A pain in the anterior part of the chest that is consistently provoked by effort and relieved by rest must be presumed to be angina, unless there is overwhelming evidence to the contrary. The pain may be provoked more easily after meals or in cold weather, but it is the relationship to effort that is of paramount importance provided that the pain develops during the exercise; a pain starting after exercise is

---

**Table C.2 Causes of chest pain**

<table>
<thead>
<tr>
<th>Acute</th>
<th>Chronic</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Central or precordial</strong></td>
<td></td>
</tr>
<tr>
<td>Common</td>
<td>Common</td>
</tr>
<tr>
<td>Myocardial infarction</td>
<td>Angina of effort</td>
</tr>
<tr>
<td>Unstable angina</td>
<td>Oesophageal spasm or reflux</td>
</tr>
<tr>
<td>Pericarditis</td>
<td>Cervical spondylosis</td>
</tr>
<tr>
<td>Dissecting aortic aneurysm</td>
<td>Prolapsed mitral cusp</td>
</tr>
<tr>
<td>Massive pulmonary embolism</td>
<td>Da Costa's syndrome</td>
</tr>
<tr>
<td>Less common</td>
<td>Less common</td>
</tr>
<tr>
<td>Upper abdominal catastrophe</td>
<td>Pulmonary hypertension</td>
</tr>
<tr>
<td>Precardial catch</td>
<td>Peptic ulcer</td>
</tr>
<tr>
<td>Acute anxiety</td>
<td>Gallbladder disease</td>
</tr>
<tr>
<td>Tracheitis</td>
<td>Chronic pancreatitis</td>
</tr>
<tr>
<td>Mycoplasma pneumonia</td>
<td>Ankylosing spondylitis</td>
</tr>
<tr>
<td>Pericardial fat necrosis</td>
<td>Tietze's disease</td>
</tr>
<tr>
<td>Lesions of sternum</td>
<td>Mediastinal tumour</td>
</tr>
<tr>
<td><strong>Lateral</strong></td>
<td></td>
</tr>
<tr>
<td>Common</td>
<td>Common</td>
</tr>
<tr>
<td>Pleurisy: infective, infarction, connective tissue disease</td>
<td>Spinal disease: infection, tumours, spondylosis</td>
</tr>
<tr>
<td>Trauma, e.g. fractured rib</td>
<td>Chronic trauma</td>
</tr>
<tr>
<td>Bronchial carcinoma</td>
<td></td>
</tr>
<tr>
<td>Less common</td>
<td>Less common</td>
</tr>
<tr>
<td>Superficial lesions, e.g. herpes zoster</td>
<td>Rib metastases</td>
</tr>
<tr>
<td>Bornholm disease</td>
<td>Aortic aneurysm</td>
</tr>
</tbody>
</table>

Very rarely, *Mondor disease*, which is phlebitis of the subcutaneous anterior thoracic veins, produces pain that is either pleuritic in type or is provoked by raising the arms. The inflamed vein may be palpable as a tender cord. Resolution occurs in a few weeks or a month or two.

**CENTRAL CHEST PAIN**

Much the most important cause of central chest pain is myocardial ischaemia. Three separate clinical syndromes are recognized – exertional or stable angina, unstable angina and myocardial infarction – although there is some overlap, at least between the first two; hence the more current term for all three is acute coronary care syndrome (ACS). ACS is then defined by the electrocardiogram (ECG) and biochemical test results into the clinical syndromes. The diagnosis of angina turns, in the great majority of cases, on an accurate history. The pain is typically symmetrical in the chest, or nearly so, being felt in the region of the sternum, or slightly to the left, and radiating laterally towards the left axilla and down the left arm. The left is involved more often than the right, but it can be bilateral or occasionally just the right. Radiation to the epigastrium, the side of the neck, lower jaw and teeth also occurs. Very occasionally, the pain is felt in the midline of the back or in one or other scapular region. The pain is described as ‘tight’, ‘gripping’ or ‘like indigestion’, or the patient may deny actual pain and describe only a feeling of pressure or of tightness. The patient may place a clenched fist placed on the sternum to indicate both the site and, presumably, the character of the sensation (Levine's sign). A pain described as ‘stabbing’ is probably not angina, but patients do not always choose their words with care, and each statement must be carefully analysed to ascertain exactly the patient's meaning. Certainly, a pain that comes in sharp jabs, lasting a second only, is unlikely to be angina, and adjectives such as ‘shooting’ and ‘stabbing’ are suspect on this account. The typical duration of an attack is only a few minutes, and a pain lasting for a much longer or shorter time than this is unlikely to be angina; there are exceptions, however, which will be discussed. Perhaps the most important aspect is the relationship to exertion. A pain in the anterior part of the chest that is consistently provoked by effort and relieved by rest must be presumed to be angina, unless there is overwhelming evidence to the contrary. The pain may be provoked more easily after meals or in cold weather, but it is the relationship to effort that is of paramount importance provided that the pain develops during the exercise; a pain starting after exercise is
not angina. The effect of sublingual glyceryl trinitrate (GTN) may be of diagnostic importance, but, again, the time relationship is important. If the pain is relieved within a minute by GTN, angina is probable; patients are, however, unfamiliar with such a rapid effect from oral medication and may claim that GTN relieves their pain, omitting the essential fact that the pain does not cease until, perhaps, half an hour after the tablet was taken. Occasionally, variants of angina are seen such as pain felt in only one of the distal sites of radiation; even ‘tennis elbow’ and ‘toothache’ may prove to be angina if the constant relationship to exertion can be elicited.

Apart from exercise, angina may be precipitated by sympathetic overactivity. This is the mechanism of angina provoked by emotion, as in the case of John Hunter, who said, some time before he died after an acrimonious Board meeting, ‘My life is in the hands of any rascal who chooses to annoy and tease me.’ Nocturnal angina is also related to sympathetic overactivity, as it has been shown to occur after a period of rapid eye movement sleep, which is associated with dreaming. Angina decubitus, provoked by lying down, is probably due to the increase in cardiac output in this posture; it is rather characteristic of syphilitic aortic valve disease. In general, angina that is easily provoked or occurs, apparently spontaneously, at rest is associated with severe disease involving all three coronary arteries (right, left circumflex and anterior descending).

In the majority of cases of angina, no abnormal physical signs are found, but two important signs should be specifically sought. A left atrial impulse may be palpable at the apex and an atrial gallop rhythm heard, particularly during an actual attack of pain. Paradoxical splitting of the second heart sound is less common and it is difficult to elicit but, if present, implies prolongation of left ventricular systole and a serious disturbance of left ventricular function. The ECG is usually normal at rest, but the typical depression of the S-T segment may be present to confirm the diagnosis of ischaemia; evidence of previous infarction may also be present or non-specific changes such as bundle-branch block. Recording the ECG during exercise to demonstrate S-T depression, and sometimes other changes, is a reliable method of confirming the diagnosis of myocardial ischaemia. Radioisotope scanning of the myocardium using, for example, thallium-201 is also helpful in delineating areas of ischaemia that develop during exercise. In the majority of patients with angina and a positive exercise test, coronary angiography will show occlusive lesions or narrowing of one or more of the coronary arteries. In a few, however, the arteries are normal or nearly so, and in such cases the most likely cause of the ischaemia is coronary artery spasm. This concept is an old one but was convincingly revived by Maseri et al. It can occur in otherwise normal arteries or in association with occlusive disease of any degree of severity.

In a large majority of cases of angina, the underlying lesion is coronary atherosclerosis. In addition, other forms of arterial disease can, occasionally, cause ischaemia. Angina is a well-recognized, albeit very rare, symptom of polyarteritis nodosa and giant-cell arteritis; smaller vessels may be involved in rheumatoid arthritis and in association with livedo reticularis. Another now rare inflammatory cause of angina is involvement of the coronary ostia in syphilitic aortitis; the attacks of pain tend to last longer and to occur rather characteristically at night, although the relationship to effort is as in other varieties of angina. Ischaemic pain is aggravated by left ventricular hypertrophy due to hypertension and, particularly, aortic valve disease. In these conditions, and in hypertrophic obstructive cardiomyopathy, the disease of the coronary arteries themselves may be trivial, and the pain is due to relative ischaemia of the hypertrophied muscle. Severe anaemia (e.g. pernicious anaemia or following gastrointestinal haemorrhage), thyrotoxicosis and tachycardia can also cause angina in patients with only minor coronary disease. All of these factors must be borne in mind, particularly when dealing with a case of angina in a premenopausal woman. Coronary atherosclerosis is rare in such patients, and angina is likely to be due to one of the precipitating factors mentioned or to premature atherosclerosis resulting from hyperlipidaemia, as in diabetes, myxoedema or hereditary hypercholesterolaemic xanthomatosis (type II hyperlipidaemia in Fredrickson’s classification).

Unstable angina is one of a number of terms used to describe cases in which prolonged cardiac pain occurs at rest without evidence of myocardial necrosis. This can be angina at rest, angina coming on with exertion but with decreasing amounts of exercise, or new onset angina causing pain on limited exercise. A variant of this condition is the so-called ‘angina inversa’, described by Prinzmetal, in which S-T elevation occurs briefly in association with ischaemic pain at rest or on exercise; this is believed to be due to severe ischaemia of a rather localized area of myocardium due to severe arterial disease, or sometimes spasm. S-T depression can also occur in similar circumstances, and it is probable that the elevation of ‘angina inversa’ is produced merely by the relationship of the site of ischaemia to the electrode position.
Myocardial infarction is nearly always due to occlusion of a coronary artery by atherosclerosis, with or without superadded thrombosis; a very rare cause is coronary embolus in association with, for example, atrial fibrillation or infective endocarditis. The pain of myocardial infarction has exactly the same character and areas of radiation as angina. It is not, however, related to exercise and typically lasts for several hours, if untreated, rather than the few minutes of an anginal attack. Myocardial infarction can sometimes be painless, especially in the elderly or in diabetics, in whom it may manifest itself as syncope, an arrhythmia or an otherwise unexplained left ventricular failure. In contrast to the paucity of physical signs in angina, it is unusual to find no abnormal physical signs in a case of myocardial infarction, provided that frequent examination is carried out since many of the signs may be very transient. Some fall in blood pressure is common but may not be detected if the previous level is unknown. Slight elevation of the jugular venous pressure is seen in many cases, and an audible or palpable atrial gallop is found even more frequently. Paradoxical splitting of the second sound may occasionally be detected, and, after a day or two, a pericardial rub may be heard.

The three most common complications are arrhythmias, cardiac failure and shock, in that order. Continuous monitoring has demonstrated that over 90 per cent of cases of myocardial infarction have some form of arrhythmia, of which ventricular extrasystoles are the most common and may presage ventricular tachycardia and fibrillation. Supraventricular arrhythmias also occur, often in association with cardiac failure. Atrioventricular block is an ominous complication, especially if it develops in a case of anterior infarction. Congestive heart failure or frank pulmonary oedema occurs from time to time, but lesser degrees of left ventricular failure are common. As a result of the consequent pulmonary venous congestion, with the probable addition of multiple alveolar collapse, mild arterial hypoxaemia is seen very frequently, the arterial partial pressure of oxygen being around 9 kPa; this often causes sufficient hyperventilation to reduce the arterial partial pressure and carbon dioxide to about 5 kPa. The term ‘cardiogenic shock’ should be reserved for cases with severe hypotension, cold, clammy skin, oliguria and clouding of consciousness.

Less common complications include rupture of the infarct, which will usually cause rapidly fatal haemoperi-cardium or, if a papillary muscle is involved, acute mitral regurgitation with pulmonary oedema; rupture of the interventricular septum causes acute right ventricular failure. Systemic embolism from a mural thrombus is not uncommon; pulmonary embolism arises most often from thrombosis of the leg veins secondary to enforced recumbency. Later sequelae are ventricular aneurysm, which may rarely calcify, Dressler’s syndrome of recurrent pericarditis and pleurisy, and the shoulder-hand syndrome, which consists of ‘frozen shoulder’ and Raynaud’s phenomenon, usually on the left.

Electrocardiography remains the most commonly used method of confirming a diagnosis of myocardial infarction, but an estimation of various serum enzymes is also valuable particularly in the (not infrequent) cases in which the electrocardiographic signs are equivocal. There are three cardinal electrocardiographic signs of myocardial infarction: a pathological Q wave, at least a third the amplitude of the R wave in the same lead and at least 0.04 seconds in duration; S-T segment elevation with an upward convexity; and T wave inversion, which may not be seen until the RS-T segment is returning to the iso-electric line, as it does during the first few weeks after the episode. The T wave may also return to, or towards, normal after some months, but the Q wave – the sign of irreversible muscle necrosis – virtually always remains indefinitely. These changes are seen in leads of which the positive terminals face the infarcted area of myocardium; in leads ‘facing’ the diametrically opposite part of the heart, reciprocal changes are seen that may occasionally be of diagnostic significance.

For descriptive purposes, infarcts are subdivided into anterior, inferior (or diaphragmatic) and ‘true’ posterior. Originally, the term ‘posterior’ was applied to the diaphragmatic surface of the heart, but now that it is possible to diagnose infarction of the small part of the left ventricle that lies posteriorly, the anatomically more correct term ‘inferior’ is preferred. To avoid confusion with the older nomenclature, infarcts at the back of the left ventricle are designated true posterior (Figs C.7–C.10).

Many intracellular enzymes are released into the circulation from the infarcted myocardium, and the rise and fall of their serum levels can be of great diagnostic value. Historically, the most commonly estimated clinically are aspartate aminotransferase (previously known as ‘glutamic-oxaloacetic transaminase’), creatine phospho kinase and lactate dehydrogenase (LDH). The first two remain elevated for 3–4 days only, but elevation of the serum LDH persists for up to 2 weeks. Much more sensitive and specific tests for myocardial damage now exist. These are various
forms of troponin which, if negative after 12 hours, are good at excluding damage. They can be positive in quite a number of other conditions, such as pulmonary embolism (PE) and atrial fibrillation (AF), and levels do rise in renal failure, so results always need to be interpreted in light of the full clinical picture.

The pain of pericarditis is, in some respects, similar to that of myocardial infarction and, as localized pericarditis is common in the latter, the differentiation may present some difficulty. Pericardial pain is felt in the sternal region and towards the left, and it may radiate to the epigastrium, neck, back, shoulders and occasionally arms. The severity varies markedly, from a mild discomfort to extreme agony, and the pain is described either as ‘stabbing’ or ‘like a knife’, or in terms very reminiscent of those used to describe the pain of myocardial ischaemia. It is aggravated by deep breathing, by coughing and by twisting movements involving the muscles of the chest wall. There is therefore some relationship with exertion, on account of the associated hyperventilation, but the aggravation by specific movements such as turning over in bed serves to distinguish it from angina. The pain is also worse in the recumbent position and is relieved by sitting up; it may also be aggravated by swallowing. The characteristic physical sign is the friction rub, heard over all or part of the precordium (see RUB, PERICARDIAL, p. 593). The third cardinal feature of pericarditis is the electrocardiogram. S-T elevation is seen in all epicardial leads, i.e. in all the leads of a conventional 12-lead record except aV_{6} (Fig. C.11). The S-T elevation is concave upwards, unlike that of myocardial infarction; pathological Q waves are, of course, absent. Later in the course of the disease, T wave inversion appears, and at this stage the differentiation from myocardial ischaemia may be difficult.
Acute pericarditis.

The diagnosis of pericarditis is incomplete unless the aetiology is determined. Common causes include virus infection (often Coxsackie B), connective-tissue disorders such as systemic lupus erythematosus (SLE), rheumatic fever, rheumatoid arthritis, bacterial infections and Dressler’s and other similar syndromes. Chronic renal failure is a well-known cause, but the most common cause of all is myocardial infarction; in that condition, the pericarditis probably contributes little to the pain.

Pericardial fat necrosis is a rare cause of pain simulating that of pericarditis. No friction rub can be heard, and the ECG is normal. A paracardiac mass may be visible in the chest radiograph.

Dissecting aneurysm causes very severe anterior chest pain that radiates to the neck, back and, later, abdomen; it rarely spreads to the arms. With dissection of the descending thoracic aorta, the pain may begin in the back. The resemblance to myocardial infarction is close and, indeed, if the dissection involves a coronary artery, infarction may in fact occur and confuse the diagnostic issue further. Important differentiating features include the absence of one or more peripheral pulses, particularly if a pulse disappears while the patient is under observation, or other evidence of arterial occlusion such as hemiparesis, blindness in one eye or haematuria. The development of aortic regurgitation, due to involvement of the aortic valve ring by the dissection, is a valuable diagnostic feature. The blood pressure is little changed compared with the fall commonly seen in myocardial infarction. The severity of the pain is of some diagnostic significance; chest pain that is hardly influenced by morphine or diamorphine may well be due to a dissecting aneurysm.

The ECG is normal unless a coronary artery is involved or pre-existing hypertensive changes are present.

Much reliance is placed on radiography in diagnosis: gross dilatation of the thoracic aorta may be seen, but this is not always easy to distinguish from unfolding unless a previous film is available. Also, the dilatation may not be very marked in the early stages. In practice, a firm diagnosis is rarely possible from the plain radiograph. If available, contrast computed tomography (CT) is the investigation of choice; otherwise, the definitive diagnosis can only be made by aortography, although even this can be misleading at times.

Another important cause of sudden anterior chest pain is a pulmonary embolism. Smaller pulmonary emboli cause pulmonary infarction (discussed below, among the causes of lateral, pleuritic chest pain). Pulmonary embolism occurs commonly in the postoperative period or during a period of enforced recumbency associated with a low cardiac output, as after myocardial infarction or in cardiac failure. In young women, oral contraceptive agents have been incriminated as the cause of the initiating venous thrombosis. The same complication may, of course, occur in pregnancy. The source of the embolism is commonly thrombosis in the veins of the leg or pelvis, but this may not have been clinically manifest except, perhaps, as a small ‘spike’ of temperature. The patient rapidly becomes severely ill with central chest pain, breathlessness, and often faintness or even loss of consciousness. Peripheral cyanosis is present, the pulse is rapid, and the blood pressure very low. Elevation of the jugular venous pressure is nearly always present; it is usually a good deal higher than in myocardial infarction causing a comparable degree of hypotension. A gallop rhythm may be heard over the right ventricle.

The ECG may show classic changes of a deep S wave in lead I, in addition to a Q wave and inverted T wave in lead III – the S1Q3T3 pattern. Other patterns include T wave inversion in V1–V4 and right axis deviation. In some cases, the ECG may be normal or reveal only a tachycardia (which is the commonest finding). The chest radiograph may show dilatation of one or both main branches of the pulmonary artery, and one lung (or part thereof) may appear unusually translucent (Westermark’s sign). The chest X-ray may, however, be normal or reveal only a pleural effusion. Ventilation–perfusion scanning may reveal multiple areas of mismatch but is often inconclusive if the patient has pre-existing cardio-pulmonary disease or an abnormal CXR. Doppler ultrasound of the legs may reveal a source for the emboli from the deep venous system. CT pulmonary angiography (CTPA) may demonstrate central or distal pulmonary emboli as well.
as giving you information about other potential causes or confounding factors relating to the chest pain. Painful angiography has largely been replaced by CTPA.

Chronic pulmonary hypertension, as in mitral stenosis or Eisenmenger’s syndrome, can produce a pain that is indistinguishable from that of angina. The cause is indeed almost certainly myocardial ischaemia as a result of the severely limited cardiac output. Severe pulmonary stenosis can cause a similar pain.

The anterior chest pain associated with prolapse of a mitral valve cusp is not usually ischaemic in origin. This lesion is not uncommon, with a prevalence of 4–5 per cent in otherwise healthy individuals, and is often associated with a mid-systolic click and late systolic murmur. The diagnosis can be confirmed by echocardiography. The pain is very variable in site, duration and severity, and has no clear-cut diagnostic features.

Pain arising from the oesophagus is felt in the midline of the chest, with radiation to the jaw, back and shoulders and, to a small extent, down the inner sides of the arms. The resemblance to angina is close, and oesophageal pain may even have some relationship to exertion, although this is never constant. The pain may be due to oesophageal spasm without any other lesion, it may occur early in the evolution of achalasia of the cardia, or it may be due to hiatus hernia with oesophageal reflux and oesophagitis. Heartburn, with a radiation of sternal pain upwards from the xiphoid, is very characteristic of the last condition. Helpful diagnostic features include the association of the pain with taking food and relief with belching; the pain of oesophageal reflux is commonly worse in the recumbent position or in other postures favouring regurgitation, such as bending forward or the cramped position of the driver of a small car, especially if wearing a tight seat belt. The demonstration of a hiatus hernia by a barium swallow is straightforward.

Other upper abdominal lesions can cause pain felt in the midline of the front of the chest. The frequency with which angina is regarded as ‘indigestion’ – even by experienced physicians suffering from this condition – bears witness to this. Catastrophes such as perforated peptic ulcer and acute pancreatitis must be remembered in the differential diagnosis of myocardial infarction. Gastric distension due to aerophagy or other causes can cause substernal discomfort, but peptic ulcer and chronic relapsing pancreatitis are less common causes of confusion; in the former, the relationship of the pain to meals is obviously important.

Finally, there is the (almost mystical) association of gallbladder disease with ischaemic heart disease. Both conditions are common, and they can certainly co-exist. Much debate has centred round whether this co-existence occurs more commonly than would have been expected by chance; many experienced clinicians have a strong impression that this is so, but none of the theories to account for it is very convincing. Suffice it to say that gallbladder pain can certainly radiate into the front of the chest and simulate angina; also, central chest pain with a constant relationship to exertion must be regarded as angina, even if gallbladder disease is present in addition.

There are numerous musculoskeletal causes of anterior chest pain. Local trauma is usually obvious, but recurrent mild trauma – for example in some particular occupations – may not be mentioned by the patient as a possible cause for the pain. The pain may be a dull ache on one or other side of the chest, rarely exactly in the midline, or it may be possible to relate it to particular movements if anterior thoracic muscles are the site of origin. Pain from spondylitis or spondylitis of the thoracic (or even the cervical) spine can be referred to the front of the chest. The distribution can occasionally simulate that of angina, and radiological proof of the spondylitis is of little diagnostic significance. Relief of the pain from wearing a cervical collar is, however, good evidence of a skeletal origin. Less obvious spinal disease may also be a cause of pain felt diffusely over the precordium or in the corresponding area on the right. Experimentally, the injection of hypertonic saline into an interspinous ligament produces pain referred to the anterior part of the corresponding dermatome with some radiation upwards and downwards. It is difficult to prove this aetiology in any particular patient, but it has been suggested that chronic mild trauma to interspinous ligaments, for example due to scoliosis, may be a cause of otherwise unexplained precordial pain.

A common cause of musculoskeletal pain includes Tietze’s disease (costochondritis). This causes pain of sudden or gradual onset in one or more upper costal cartilages. The pain is worse on coughing or deep breathing, and the affected cartilage is swollen and tender. Xiphoidalgia is a similar condition involving the xiphisternum that is possibly a variant of Tietze’s disease, or more likely due to recurrent mild trauma. Ankylosing spondylitis can cause a diffuse pain in the anterior chest wall often associated with local tenderness over the sternum and costal cartilages. Pain arising in the sternum itself may be due to myelomatosis, metastases, ankylosing spondylitis,
osteomyelitis or fracture. The sudden, sharp pain at or near the cardiac apex (known as precordial catch) is probably quite common, although rare as a spontaneous complaint. There is rarely any diagnostic confusion with other causes of chest pain as the history is quite characteristic. The pain occurs most often when the subject is seated and lasts for a few minutes only. It can be relieved by a single, painful, deep inspiration. Respiratory disease most often, if at all, causes pain in one or other side of the chest, but tracheitis should be mentioned as a cause of upper sternal pain. It is aggravated by the hyperventilation of exercise and may thus, very occasionally, have to be distinguished from angina. Mycoplasma pneumonia, unlike other varieties of pneumonia, is never associated with pleurisy, but may cause a substernal pain aggravated by coughing. Dyspnoea itself – especially if it is associated with obstruction of the airways – may be described as ‘tightness in the chest’, and an injudicious series of leading questions may persuade the patient that he or she has a ‘tight pain’ across their chest and lead to an erroneous diagnosis of angina.

Two psychological conditions are important in the differential diagnosis of anterior chest pain. The terrifying panic of an attack of acute anxiety can be confused with a myocardial infarction or other intrathoracic vascular catastrophe. The patient, very often a woman, complains of dizziness, palpitations, dyspnoea and precordial oppression or pain. Angor animi – the fear of impending death – is a prominent feature, much more so than in genuine myocardial ischaemia, and further exacerbates the anxiety. The circumstances in which the attack occurs, and the complete absence of objective evidence of organic disease, are important distinguishing features, and doubt about the nature of the condition should rarely persist for long.

Chronic anxiety is the most common underlying disorder in Da Costa’s syndrome; the non-committal eponymous title is preferred to the numerous other terms, such as ‘effort syndrome’, ‘cardiac neurosis’ and ‘disordered action of the heart’, which have been used (with no great semantic accuracy) to describe this condition. Apart from the pain, the features of the syndrome are dyspnoea, palpitation, fatigue and dizziness. The pain is most commonly felt in the left submammary region, although it may be nearer the midline, or indeed anywhere on the left side of the chest, even radiating into the left arm. In character it is sharp and stabbing, with occasional momentary twinges superimposed on a dull ache that persists for many hours at a time. It often occurs after – but rarely during – exertion; this important point may be elicited only after careful questioning. In summary, it differs from angina in almost all respects except in the rare, difficult case in which it has a more constricting quality and is felt near the left border of the sternum. The mechanism of the pain is unknown, but it seems likely that, in some cases at least, the original cause was a minor musculoskeletal abnormality. The pain convinces patients that their heart is diseased and is perpetuated by the anxiety engendered by this conviction.

LATERAL CHEST PAIN
Almost all tissues of the lateral chest wall – the pleura, muscles, ribs and intercostal nerves – can be the site of painful lesions. Frequently, the pain is closely related to respiration, as exemplified most clearly in pleurisy. The visceral pleura is insensitive, but the parietal pleura is plentifully supplied with pain fibres from the intercostal nerves. The pain is felt, therefore, in the cutaneous areas supplied by these nerves, which, it is important to remember, include a large area of the anterior abdominal wall. Apart from inflammation of the pleura itself, there is some evidence that spasm of the intercostal muscles may be a factor in producing the pain. The pain is characteristically sharp, superficial and of any degree of severity, and is aggravated by deep breathing and by coughing. Inspiration is abruptly halted by the pain so that respiration is often very shallow. Holding the breath in expiration will usually relieve the pain completely, and a change in the patient’s posture in bed can produce considerable relief or exacerbation. The only physical sign of pleurisy, in the absence of effusion, is the pleural friction rub – a characteristic creaking sound present during inspiration and expiration. The sound is difficult to describe, but easy to recognize with a little experience. The pain is not at all closely related to the rub as one may be present without the other; a rub is not essential for the diagnosis of pleurisy. The pain of diaphragmatic pleurisy is typically referred to the shoulder; this is a common feature of pleurisy due to subdiaphragmatic lesions such as liver abscess or subphrenic abscess.

Pleurisy may be due to pulmonary infections such as lobar pneumonia or tuberculosis, to vascular lesions such as pulmonary infarction, or to connective-tissue disorders such as SLE. In many of these conditions, the clinical picture is modified by the development of an effusion, which usually results in the disappearance of the pain and the rub.

A pain identical to that of pleurisy is felt in epidemic pleurodynia or Bornholm disease, due to group B Coxsackie viruses. The pain is the presenting symptom
and may be extremely severe; fever quickly develops, and headache and malaise are common. Recovery is usually rapid, but relapses are frequent and may continue for several weeks. Trichinosis, involving the intercostal muscles, can also produce pleuritic pain; the diagnosis would be supported by finding periorbital or generalized oedema, and by eosinophilia in the peripheral blood. Pain arising from the capsule of the spleen is also related to respiration; it is commonly due to splenic infarction, and may be accompanied by a friction rub so that the resemblance to pleurisy is very close. Splenic pain is not uncommon in Hodgkin’s disease and in other similar conditions.

The pain of spontaneous pneumothorax is usually abrupt in onset (Fig. C.12). Some patients, however, complain only of a dull ache or a sense of tightness or often may have no pain at all. The typical physical signs are a diminution in movement and in breath sounds on the affected side, often with a hyperresonant percussion note and – especially if the pneumothorax is under tension (medical emergency) – deviation of the trachea towards the normal side. Dissection of the air into the mediastinum may cause central chest pain, and the patient may notice a ‘crunching’ sound over the heart that is also audible on auscultation. The diagnosis of spontaneous pneumothorax can be confirmed by radiography, but careful study may be needed if the pneumothorax is shallow; a film taken in expiration will show the lesion more clearly.

Involvement of the intercostal nerves in many pathological processes can cause pain in the corresponding areas of the chest wall. Spinal disease has already been mentioned as a cause of referred pain in the anterolateral regions of the chest. Direct pressure on nerves may occur in fracture of the thoracic spine, from malignant metastases in that region, or in tuberculosis of the spine. Spondylosis with disc protrusion is not common as a cause of nerve root compression in the thoracic spine. Neurofibromatosis may affect the thoracic nerve roots, but does not often cause pain.

Aortic aneurysm is not a common lesion, although in former years it was an important cause of chest pain. Aneurysm of the ascending aorta may cause chest pain by eroding the sternum, but much more often causes no symptoms at all. Aneurysm of the arch and descending thoracic aorta can cause very severe radiating pain by vertebral erosion and pressure on nerve roots. Other symptoms and signs result from pressure on mediastinal structures. Thus, pressure on the left recurrent laryngeal nerve will cause paralysis of the left vocal cord, while cough and stridor may follow pressure on the trachea, and dysphagia results from pressure on the oesophagus. Pressure on the left main bronchus may cause collapse of the left lung with subsequent infection; a tracheal tug is a well-known physical sign of aneurysm depressing the left main bronchus.

Primary or secondary intrathoracic malignant disease may cause pain in various ways. Direct invasion of the pleura by a bronchial carcinoma can cause pleurisy, often with effusion; more often, pleural pain occurs as a result of infection in the lung distal to a blocked bronchus. Primary tumours of the pleura (e.g. mesothelioma) cause a range of pains from mild ache to severe and constant pain directly. Apart from the pleura, the ribs and intercostal nerves may be involved by tumour with the production of severe pain. Metastases in the thoracic spine have been mentioned as a cause of intercostal pain; secondary deposits in the ribs can also be extremely painful. Rarely, tumours in the mediastinum can apparently cause a poorly localized central chest pain without other pressure symptoms; the mechanism is not known.

**CHEST, TENDERNESS IN**

**Alex West**

Chest pain is one of the most common symptoms given for seeking medical advice. Because there is no clear relationship between the intensity of...
discomfort and aetiology, all complaints of chest pain must be considered carefully (see also CHEST PAIN, p. 88). This section deals exclusively with tenderness, which is, in fact, rarely complained of in the absence of pain.

Tenderness in the chest, an ache or discomfort, perhaps with increased sensitivity and often accompanied by pain, can be difficult for the patient to describe. It is best classified according to the situation or character of the responsible lesion. Pains referred from visceral lesions may sometimes be associated with local tenderness in the chest wall. The parietal pleura is exquisitely sensitive to painful stimuli which to the patient may seem like their chest wall but the same pain is not replicated on palpitation, and unpleasant sensations may also arise from lesions in the tracheobronchial tree.

Lesions of the chest wall
- Inflammation of the skin and underlying tissues including the breasts
- Intercostal myositis
- Myalgia
- Inflammation of the ribs and sternum
- Primary and secondary malignancies
- Intercostal neuritis and neuralgia
- Injury of the intercostal nerves
- Ankylosing spondylitis
- Herpes zoster
- Tietze's syndrome

Lesions of thoracic and abdominal viscera
- Lungs
- Heart and aorta
- Diaphragm
- Stomach and oesophagus
- Liver/gallbladder

LESIONS OF THE CHEST WALL
Discomfort in the chest wall can result from respiratory diseases, as well as from primary musculoskeletal lesions. Patients with chronic cough, dyspnoea or asthma subject to chest tightness often complain of anterolateral chest wall tenderness.

Tenderness is always present in superficial inflammatory lesions of the chest wall, such as bruises, burns, cuts, mastitis and superficial infections, the diagnosis of which will usually be evident on examination.

Pain will be the chief complaint in the so-called intercostal myositis that occurs after injury or strain of an intercostal muscle, the affected muscle being tender to deep pressure. The condition is also known as intercostals myalgia or pleurodynia. It is distinguished from pleurisy by the absence of a pleural friction rub. Similar but more transient pain with a variable degree of tenderness may accompany the stitch to which some athletes are prone.

The acute pain of Bornholm disease – epidemic myalgia due to Coxsackie virus B infection – may be accompanied by hyperaesthesia of skin, but less often by muscle tenderness. The myalgia of Phlebotomus (sandfly) fever and dengue may also be accompanied by tenderness, often mild.

Tenderness of the breasts in the absence of mastitis is a common occurrence at or just before the menstrual periods and with high-dosage oestrogen therapy. Gynaecomastia in males, whatever the cause, is accompanied by tenderness of the breasts. It is not uncommon in males who are chronic alcoholics and have cirrhosis of the liver.

Tenderness in the chest may result from disease or injury of a rib or the sternum, when it will be localized to the injured spot; fracture, inflammation, tuberculosis or new growth may be the immediate cause. If a fracture is present, X-rays may show the lesion, or crepitus between the fragments on movement may be obtainable. Sternal or costal ostelitis or periostitis may follow injury and may also occur in such diseases as typhoid or paratyphoid fever and tuberculosis. The local signs of inflammation (pain, redness, heat and swelling) will usually but not invariably be present. The chest wall may be invaded by local extension of a peripheral primary bronchial carcinoma or secondary tumour. Tenderness in the chest due to malignancy in a rib or in the sternum, such as multiple myeloma, sarcoma or a secondary deposit from carcinoma, is generally a late occurrence, the existence of malignant disease elsewhere having usually been established. Tenderness of the ribs and sternum occurs in certain blood diseases such as leukaemia. Diagnosis depends on examination of the blood and marrow biopsy.

Tenderness over the sternum and ribs also occurs as part of the clinical picture of ankylosing spondylitis. In this disease, the sternomanubrial and sternoclavicular joints may become acutely swollen and tender, causing considerable discomfort.

The particularly tender spots in the course of an intercostal nerve are three in number, corresponding to the points at which the posterior, the lateral cutaneous and the anterior cutaneous branches are given off, near the spinal column, in the mid-axillary line and at the sternal margin, respectively. Such tenderness
may be marked in so-called intercostal neuritis, when some intrathoracic disorder such as pneumonia or pleurisy is present and in cases of pressure on an intercostal nerve, as for example by abscess about the spinal column, aneurysm of the descending aorta or malignancy invading the spinal canal. Local tenderness may more commonly result from external pressure by, for instance, the buckle of the braces or some tool carried in a breast pocket, a simple detail but one not infrequently overlooked.

Pain and tenderness along an intercostal nerve are common in herpes zoster and may be present before, during and after the appearance of the characteristic rash. Tenderness can often be elicited at the three spots mentioned above; it is particularly when it occurs past middle-age that herpes may be followed by a long period of pain and tenderness along the course of the affected nerve. The rash, once seen, can hardly be mistaken; to anticipate it on the type and site of the pain is a diagnostic tour de force. Similar pain and tenderness may follow thoracotomy, and occasional patients experience intractable postoperative discomfort.

Tietze’s syndrome is an unexplained disorder in which pain and swelling are found in the upper costochondral junctions of the anterior chest wall. Biopsies of the costal cartridges show nothing characteristically abnormal. One or more costal cartilages, usually on one side only, and most often the second and third, may be affected. Spontaneous remission occurs in weeks, months or occasionally years.

**LESIONS OF THE UNDERLYING VISCERA**

Tenderness in the chest may sometimes be a symptom of disease in the thoracic or abdominal viscera. The tenderness is as a rule superficial, confined to the skin and subjacent areolar and fatty tissues. Tactile hyperaesthesia or the production of unpleasant sensations or pain by the lightest touch may occur in neuralgia, in neuroses, following thoracotomy or in cases of referred pain. A similar hyperaesthesia for cold or less often for heat sometimes occurs in the chest of tabetic patients. Hyperalgesia, where a normally painless stimulus becomes transformed into an acutely painful sensation, may be regarded as a form of ‘tenderness’ in the chest. This occurs in patients suffering from anxiety states, often with added depression. Furthermore, perversions of sensation sometimes occur in organic nervous diseases, such as syringomyelia or tabes.

Tenderness of the chest may occur in pleurisy. The tenderness is as a rule deeply seated, and not in the skin and subcutaneous tissues. However, chest wall pain can mimic pleurisy, and conditions in the chest wall can cause confusion. Tenderness due to strain or tearing of the thoracic muscles can be severe and painful, may be exacerbated by coughing and may be confused with pleurisy.

The sternum may be tender as the result of mediastinal inflammation, tumour or aneurysm. The diagnosis in these cases is made by physical and X-ray examination. Tenderness with pain over the precordium may occur in pericarditis, accompanied usually by a pericardial rub. It may be so severe as to preclude percussion or even the application of a stethoscope. Similar pain and tenderness may also be found in the epigastrium and upper costal angles.

Chest tenderness is sometimes found in cases of acute or chronic disease of the lungs, particularly tuberculosis. The tenderness may be either superficial or deep. It is generally felt most about the region of the apices of the lungs, the curve of the shoulder or the scapula. Similar tenderness is met with occasionally in acute bronchitis, or in chronic bronchitis and emphysema. Tenderness along the lower chest wall anteriorly may be found after vigorous coughing, probably from trauma in the soft tissues, the muscles particularly. A rib may be fractured by vigorous coughing.

Direct tenderness about the precordium is almost never due to heart disease. It is more generally associated with cardiac neurosis than with organic heart disease. Tenderness at the area of the apex beat is common in the Da Costa’s syndrome (‘soldier’s heart’ or ‘neurocirculatory asthenia’), a nervous condition in which there is no cardiac abnormality. The tenderness, which may be extreme, felt by some patients with heart disease, such as mitral stenosis, at the cardiac apex is due to anxiety rather than to an organic lesion.

Tenderness in the right side of the chest near the costal margin is not rare in diseases of the liver and gallbladder corresponding to the cutaneous distribution of the D7, D8 and D9 nerves; for the most part, however, the pain and tenderness are in the epigastrium and the right hypochondrium. The right phrenic nerve (C3–C5) sends branches to the liver and gallbladder, so that tenderness and pain may also be felt in the right shoulder, as in the case of disorders of the diaphragm. It is particularly in cases of gallstone or biliary colic that these areas of tenderness are likely to be found. In patients with hepatic abscess, the spread of inflammation to the chest wall may give rise directly to pain and tenderness.
CHEYNE–STOKES RESPIRATION

Alex West

This well-known abnormality of respiration, described independently by John Cheyne in 1818 and William Stokes in 1846, is probably referred to in the Hippocratic writings in an account of a patient whose breathing was ‘like that of a man recollecting himself, and rare and large’.

Cheyne–Stokes respiration – the most common form of periodic breathing – consists of alternating periods of apnoea and hyperventilation, beginning with hardly perceptible movements, gradually increasing until the tidal volume is much above normal, and then dying away to end in apnoea (Fig. C.13). The apnoeic period lasts for 10–30 seconds or more, and the hyperpnoeic phase comprises 30 or more breaths and usually lasts between 1 and 3 minutes. The condition is obvious to the experienced observer, but an untrained person will often describe the hyperpnoeic phase as ‘breathlessness’. The patient may be unaware of the breathing abnormality. As Cheyne–Stokes breathing is accentuated during sleep, the hyperpnoea may disturb the patient’s sleep, and the symptoms may be confused with those of paroxysmal nocturnal dyspnoea due to cardiac failure.

The mechanism responsible for Cheyne–Stokes breathing is complex. In health, there is an oscillating balance between changes in arterial blood gas tensions and respiratory drive. The system is controlled by peripheral and central chemoreceptors, the rate of response of which is dependent upon the time it takes the circulation to carry arterial blood from the lungs to the carotid bodies and to the brain. The effect of these functional changes is most apparent when the predominant regulator for breathing is by the chemical control system that occurs in non-rapid eye movement (REM) sleep. Thus, in stages 1 and 2 of non-REM sleep, periodic breathing is normal and is initiated by a change in the homeostatic set point for ventilation induced by the onset of sleep. Physiological periodic breathing is accentuated at altitude.

Pathologically, Cheyne–Stokes respiration – which is an extreme form of periodic breathing – occurs almost exclusively during non-REM sleep and typically ceases during REM sleep. Periodic breathing commonly signifies cerebral or cardiac disease, and is most common following a stroke or in cases of left ventricular failure. At least three mechanisms have been postulated. First, in patients with an increased respiratory drive due to chronic hypoxaemia, a further fall in arterial oxygen gas tension (PaO₂) at the onset of sleep may result in periodic breathing by a mechanism analogous to that occurring in healthy subjects at altitude. Second, there is an inevitable circulatory delay in arterial carbon dioxide tension (PaCO₂) changes produced by altered ventilation and the detection of blood gas changes by the central chemoreceptors. The lung-to-brain circulation time may be prolonged in certain cardiac or cerebrovascular diseases. Third, the chemoreceptors may be over-responsive to changes in PaCO₂ due to a loss of normal inhibitory influences on the metabolic control system, such as may occur in bilateral pyramidal tract destruction.

Thus, in normal subjects, voluntary hyperventilation with air will lead to a short period of apnoea followed by a few cycles of Cheyne–Stokes breathing. It is therefore possible to reduce the PaCO₂ to such a level that even a healthy respiratory centre fails to discharge normally; this does not occur after hyperventilation with 5 per cent carbon dioxide. The slow decline in arterial oxygen saturation and rise in carbon dioxide during the apnoea begins to stimulate the respiratory centre, and respiration is resumed either normally or leading to a second fall in PaCO₂ with repetition of the cycle. The changes in the blood gases during the cycle are shown in Figure C.14.

Cheyne–Stokes respiration occurs in normal subjects not only after hyperventilation but also at high altitude, where the hypoxic stimulus to respiration reduces the PaCO₂. It may also be seen in apparently healthy elderly subjects during sleep; it is difficult, however, to exclude minor degrees of cardiac or cerebrovascular disease causing depression of the respiratory centre.

**Figure C.13** Spirogram from a patient with severe cerebral vascular disease. Two cycles of Cheyne–Stokes breathing are shown over a period of 143 seconds.

**Figure C.14** Diagram of changes in tidal volume, arterial oxygen saturation (SaO₂) and partial pressure of carbon dioxide in arterial blood (PaCO₂) over two cycles of Cheyne–Stokes breathing.
CHOREA

in such cases. In clinical practice, the most common cause is left ventricular failure. Periodic breathing occurs especially in patients with degenerative arterial disease in whom the blood supply to the brainstem may be reduced as a result of the low cardiac output and local arterial disease. Cheyne–Stokes respiration is commonly regarded as indicating a poor prognosis in left ventricular failure, but it may disappear with treatment for the failure, and may rarely persist for many months in patients in whom the other symptoms and signs of failure are unimpressive.

Bronchopneumonia or other respiratory infections may also precipitate Cheyne–Stokes breathing in the elderly. However, it must be realized that in chronic respiratory failure, in which a raised rather than a lowered PaCO2 is the rule, Cheyne–Stokes respiration does not occur. Occasionally, there may be a few cycles in the recovery period following a Stokes–Adams attack. Respiration continues during the period of circulatory arrest, and the first blood which enters the cerebral circulation after cardiac action is resumed contains very little carbon dioxide. The sensitivity of the respiratory centre is reduced by hypoxia during the circulatory arrest, and this combines with the hypocapnia to cause Cheyne–Stokes breathing. Rarely, Cheyne–Stokes breathing is complicated by cardiac arrhythmias, including junctional rhythm and atrioventricular block, which occur intermittently in phase with the respiratory arrhythmia; the mechanism of this is unknown.

Primary depression of the respiratory centre in the absence of much change in the PaCO2 can also cause Cheyne–Stokes respiration. Thus, it occurs in many diseases of the central nervous system. These include cerebral vascular disease with or without haemorrhage or thrombosis, cerebral tumours (especially those involving the brainstem) and severe head injuries. Cheyne–Stokes respiration is always more prominent during sleep, and it can be precipitated by the administration of narcotic hypnotic drugs such as morphine, or occasionally benzodiazepines. It is also seen quite often in uraemia but is probably not due to the renal failure per se. Hyperventilation in renal failure is caused by acidosis, the effect of which persists despite the fall in PaCO2. Left ventricular failure resulting from renal hypertension may be responsible for Cheyne–Stokes breathing in this situation, although it may occur in patients whose blood pressure is normal.

Cheyne–Stokes respiration may be confused with other periodic breathing patterns that typically show a shorter, less symmetrical, and regular contour. In pontine brainstem lesions or if the intracranial pressure is raised abruptly, short clusters of hyperpnoeic breathing may be interrupted by abrupt spasms of apnoea (Biot’s breathing). Ataxic respiration may be seen with medullary lesions, which can provoke a grossly irregular breathing pattern. Respiratory apraxis is recognized by a monotonously regular pattern of breathing that cannot be modified, and is seen in the ‘locked-in’ syndrome when subjects suffer bilateral pyramidal lesions. Central neurogenic hyperventilation is seen occasionally in midbrain lesions.

CHOREA

David Werring & Mark Kinirons

Chorea is derived from the Greek word for ‘dance’ and refers to involuntary, irregular, purposeless, non-rhythmic movements of a rapid and jerky type that flow from one part of the body to another. This is in contrast to the slow, writhing movements of athetosis. Choreiform movements may be simple or elaborate, and typically flit unpredictably from one region to another. They are purposeless, but attempts may be made to incorporate them into functional movements by the patient. The movements may affect the face, causing grimacing; there may be movements of the tongue, and also peculiar grunting sounds on respiration. The involuntary movements, which may affect any of the muscles of the body, are made worse by attempted voluntary movement, excitement and maintenance of posture. Normal volitional movements are possible but are frequently interrupted by the chorea. If the patient is asked to grip the examiner’s hand, the force will be felt to vary; this physical sign is termed motor impersistence or ‘milkmaid’s grip’.

It is frequently difficult to dissociate chorea from the slower, writhing movements of athetosis, and the term ‘choreoathetosis’ is sometimes used to describe involuntary movements that have characteristics of both. It is also important to differentiate chorea from myoclonus, the main distinguishing features being that myoclonus is much more rapid and does not flow from one muscle to another like chorea. There are many causes of chorea, some of which are listed in Box C.3. A few of the more important causes are discussed further below.

Sydenham’s chorea is a disease that follows rheumatic fever due to group A beta-haemolytic streptococci. The mechanism has recently been postulated to involve antibodies, generated as part of the response to infection, that cross-react to basal ganglia structures. Chorea gravidarum occurs during pregnancy and in patients taking oral contraceptives, and it may follow an earlier episode of rheumatic fever.
of Huntington’s disease, it carries a more benign prognosis. Some cases previously ascribed to this category may in fact be due to cerebral small-vessel disease affecting the basal ganglia. **Congenital chorea** is sometimes seen in association with congenital hemiplegia and diplegia but is less common than congenital athetosis. Choreoathetosis in the young may be seen in a number of rare inherited degenerative metabolic disorders (see Box C.3). **Hemichorea** is often due to a structural lesion (vascular damage or neoplasm). A strikingly violent movement disorder related to, but distinct from, chorea is *hemiballismus*. It is seen most often in elderly patients with diabetes and hypertension due to small vascular lesions (lacunar infarction), classically within the contra-lateral subthalamic nucleus. The syndrome begins abruptly, but the movements tend to decrease in magnitude, and the patient may be left only with minor irregular flexion or extension movements of the wrists and fingers.

**CLONUS**

David Werring

Clonus is a series of rhythmic, monophasic (i.e. unidirectional) contractions and relaxations of a group of muscles. This is in contrast to tremors, which are always bidirectional. Clonus is a component of upper motor neurone syndrome.

**CLONUS IN UPPER MOTOR NEURONE LESIONS**

Clonus is one of the positive (excess) motor phenomena that characterize upper motor neurone (UMN) syndrome. The UMN syndrome results from damage to descending pathways to the lower motor neurones (anterior horn cells) in the spinal cord, especially the corticospinal tract. Damage to these pathways – which can be at a spinal or a cerebral level – interrupts the supraspinal control of spinal reflexes, causing the positive phenomena of clonus, spasticity, hyperreflexia and reflex spread. These features often, but not always, occur together.

Clonus is a series of rhythmic, involuntary muscle contractions occurring at a frequency of 5–7 Hz in response to an abruptly applied and sustained stretch stimulus; it is frequently seen at the ankle but may also be elicited at the patella. It depends upon voluntary relaxation of the muscles, the integrity of the spinal stretch reflex mechanisms, sustained hyperexcitability of the alpha and gamma motor neurones, and synchronization of the contraction–relaxation cycle of the muscle spindles. Several beats of clonus (fewer than five or six) may be elicited normally at
the ankle by abrupt dorsiflexion of the foot. More numerous beats, or sustained clonus, imply an upper motor neurone lesion above the level of the first sacral segment and usually occur in association with an extensor plantar reflex. When the upper motor neurone lesion is above the third and fourth lumbar segments, patellar clonus may also be elicited.

CONFUSION

Mark Kinirons

Confusion is the term used to indicate that a subject is temporarily unable to think in a clear and logical fashion. The value of the term in diagnosis is, however, much diminished by imprecise definition. In a confusional state, some or all of the following features are found: an impairment of concentration and attention span with an inability to shift attention to new external stimuli; memory impairments; disorientation in time, space or person; speech that is rambling, irrelevant or incoherent; and an impaired ability to properly grasp the meaning or significance of surrounding events. These features are extremely common and are found in a wide range of disorders that fall into the following main categories: acute confusional states or delirium; chronic confusional states (chronic organic reactions or the dementias); the functional psychoses; dissociative states; and other neurotic conditions. The term should be considered as a convenient clinical description of a certain qualitative change in consciousness.

ACUTE CONFUSIONAL STATES (DELIRIUM)

The term 'acute confusional state' and 'acute brain syndrome' should be considered as synonymous with 'delirium'. The speech is incoherent and rambling, indicating disordered thinking. Questions have to be repeated because the subject's attention is poor. There is an altered level of consciousness; this may be reduced in approximately 30 per cent of patients, but is increased in 70 per cent. The onset is usually rapid, and the course often fluctuating. There is a disturbance of the sleep–wake cycle. Fearfulness is common, and many patients experience visual hallucinations. Diagnosis is aided by Confusion Assessment Method (CAM) questionnaire.

The range of possible causes of delirium is large. These causes frequently lie outside the nervous system and include infectious, metabolic and toxic states due to alcohol and drugs (pharmaceutical and recreational). A careful history, complete physical examination and laboratory investigations suggested by the features of the case will generally be sufficient to identify the cause (Box C.4).

### BOX C.4 CAUSES OF ACUTE STATES OF CONFUSION (DELIRIUM)

<table>
<thead>
<tr>
<th>Common Causes of Acute States of Confusion (Delirium)</th>
</tr>
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<tbody>
<tr>
<td>Common:</td>
</tr>
<tr>
<td>• Hypoxia (due to cardiac and respiratory disorders)</td>
</tr>
<tr>
<td>• Metabolic disorders</td>
</tr>
<tr>
<td>• Systemic infections</td>
</tr>
<tr>
<td>• Cerebral infections</td>
</tr>
<tr>
<td>• Non-infective cerebral causes</td>
</tr>
<tr>
<td>More common:</td>
</tr>
<tr>
<td>• Hypothermia</td>
</tr>
<tr>
<td>• Seizure (status)</td>
</tr>
<tr>
<td>• Post-ictal state</td>
</tr>
<tr>
<td>Less common:</td>
</tr>
<tr>
<td>• Focal lesions of right parietal lobe</td>
</tr>
<tr>
<td>• Porphyria</td>
</tr>
<tr>
<td>• Hypothyroidism</td>
</tr>
<tr>
<td>• Hypoparathyroidism</td>
</tr>
<tr>
<td>• Hypopituitarism</td>
</tr>
<tr>
<td>• Cushing's disease</td>
</tr>
</tbody>
</table>

CHRONIC CONFUSIONAL STATES (CHRONIC ORGANIC REACTIONS)

(See also AMNESIA p. 21; MEMORY, DISORDERS OF, p. 400)

These are sometimes broadly defined as 'the dementias'. Dementia is defined as impairment in two or more areas of higher cortical function with a change in function of daily activities and the emergence of new behavioural symptoms, with the consequent social impact. Examples include forgetting money, misplacing items and getting lost on the way home. Alzheimer's disease accounts for approximately 60 per cent of all cases of dementia; the remainder are due to multi-infarct dementia (30 per cent) and miscellaneous causes (5–10 per cent) (Box C.5 and Fig. C.15).

The most common presenting feature is impairment in short- and long-term memory, often accompanied by other features of higher cortical dysfunction such as impairment of abstract thinking and judgement. Relatives may notice a gradual personality change and a depressive, irritable, disinhibited or euphoric mood disturbance. The memory loss initially may be mild and most marked for recent events. The person forgets names, loses objects round the house, or
experiences distressing confusion in unfamiliar surroundings due to loss of spatial memory. As the condition progresses, new information is not retained, and the person may become forgetful enough to become a danger to themselves by leaving tasks undone and taps and switches left on. Impairment in abstract thinking is suggested when the patient cannot cope with new tasks, thinking becomes more literal and concrete, and the repertoire of conversation is narrowed. There may be impaired judgement, the person becoming disinhibited with inappropriate behaviour in social situations. Marked disinhibition can be a feature of frontal lobe impairment. Other neurological deficits including aphasias and apraxias are common, as are changes in mood, usually to depression and irritability.

Alzheimer’s-type dementia is diagnosed by the clinical findings and the exclusion of other causes of a progressive dementia. In multi-infarct dementia, there may be hypertension and a stepwise progression of impairment with a history of transient ischaemic episodes. Computed tomography (CT) scanning may show old infarcts, whereas in Alzheimer’s dementia the CT scan may be either normal or show selective cerebral atrophy of the temporal lobes and ventricular enlargement. A variant of Alzheimer’s disease, known as posterior cortical atrophy presents with predominantly visual impairment and inability to conduct tasks due to more selective atrophy of the visual cortex.

Lewy body dementia is increasingly recognized as a cause of dementia due to an excess deposition of Lewy bodies in the brain’s subcortical structures. This leads to presenting symptoms differing from those of Alzheimer’s dementia or multi-infarct dementia. Typically, patients have psychomotor retardation, prominent visual hallucinations, a fluctuating level of consciousness (from day to day) and exaggerated sensitivity to major tranquillizers. The disease is incurable and progressive.

Metabolic disorders giving rise to chronic confusional states include vitamin B₁₂, folate and thiamine deficiencies. The latter (Wernicke–Korsakoff syndrome) is commonly associated with chronic alcoholism but may be due to gastric carcinoma or malabsorption. Cerebral anoxia due to anaemia or respiratory or cardiovascular disease also causes symptoms of confusion, although these latter causes are more likely to present with acute confusion.

Brain degenerative conditions giving rise to dementia include Huntington’s disease, which is an autosomal dominant condition usually presenting in the age
range of 35–45 years, with involuntary movements and cognitive changes. A proportion of patients with multiple sclerosis develop a steadily progressive illness with a severe impairment of higher cortical function. There are usually other signs and symptoms disseminated in time and place. In normal-pressure hydrocephalus, a dementia is typically accompanied by apraxic gait and urinary incontinence that occurs early in the dementing illness, in contrast to Alzheimer’s disease, when it is late. Pick’s disease, which is the most common of a family of frontotemporal dementing syndromes, is a rare condition that usually develops in older patients, typically with a marked personality change consistent with frontal lobe impairment and a speech disorder that progresses to mutism. This is due to a selective loss of nerve tracts to the frontal cortex. Recent advances have identified gene defects leading to frontotemporal dementia in certain families. Signs of cognitive impairment are found in Parkinson’s disease, especially in older patients who have had the condition for a long time. Binswanger’s leukoencephalopathy is a very rare progressive degenerative disease with signs on CT scan of focal white matter sclerosis, which are characteristic of an inability to relax the hand grip, and hypertention.

Central nervous system infections that give rise to features of dementia include viral encephalitis. The human immunodeficiency virus is neurotropic as well as lymphotropic, and people with AIDS may develop a confusional state with fever leading to a state of lethargy, apathy and ataxia due to a subacute encephalopathy. Tertiary syphilis is now rare in the West, but it is still an important, treatable, cause of dementia worldwide. Creutzfeldt–Jakob spongiform encephalopathy, caused by a prion protein, is also quite rare and has features including limb weakness with spasticity, myoclonus cerebellar degeneration and other movement disturbances. Electroencephalographic (EEG) changes are common. Variant Creutzfeldt–Jakob disease is also very rare, being due to the ingestion of prion protein particles in the diet or their acquisition through surgery. Cases have been reported worldwide.

Psychiatric illness can present as confusion. Schizophrenia, which may remain undiagnosed until relatively late in life, can present with a degree of intellectual deterioration, impaired volition, disordered thinking and confused speech. A detailed history may reveal a family history or earlier episodes of psychotic illness, but memory loss is not a feature. Pseudodementia is a condition of patients with major depressive illness that presents a dementing-like picture of poor memory, impoverished thinking and reduced intellectual abilities, and it may mimic an organic dementia. This condition is diagnosed by careful history-taking from the patient and relatives, and it should be borne in mind that in every case of ‘dementia’, the depression can be cured by treatment. Very rarely, a factitious disorder may mimic acute or chronic confusional states, but careful assessment will usually reveal inconsistencies in the patient’s account of symptoms and their performance in tests of higher cerebral function. Epileptic twilight states and automatisms are characterized by an abrupt onset and ending, a duration of hours or rarely days, and the occurrence of apparently purposeless acts. The subject has impaired consciousness. A history of seizures and EEG evidence of epileptic activity should be sought and would be essential evidence for a patient facing prosecution for offences claimed to have been perpetrated during a state of epileptic automatism.

CONSCIOUSNESS, DISORDERS OF

Mark Kinirons

Consciousness is the state of awareness of the self and the environment when provided with adequate stimuli. Patients who are in normal wakefulness will be fully responsive to stimuli and will display correct behaviour and speech in response to these stimuli. Physiological changes in consciousness occur during sleep, when the patient may be easily aroused by external stimuli. Pathological impairment of consciousness occurs in states of injury, disease or intoxication.

A useful measure to assess level of consciousness is the Glasgow Coma Scale (Fig. C.16). It is based on observation of levels of response to graded stimuli in three modes: eye-opening, motor function and verbal utterance. In the first, spontaneous eye-opening and blinking score the highest grade, eye-opening in response to voice the next highest, eye-opening in response to painful stimuli the next, and finally no response of eye-opening to stimuli the lowest. The highest motor response is voluntary movement to command, followed by a localizing response to painful stimuli, withdrawal from painful stimuli, a pathological flexor response, sometimes called decorticate, a pathological extensor response, sometimes called decerebrate, and finally no response at all to pain. In terms of verbal response, the highest level is orientated conversation, the next disoriented conversation, followed by the use of occasional recognizable words, no recognizable words but groaning in response to pain, and finally no response of a verbal nature. By allocating a score in each of these categories, the
level of consciousness of the individual patient may be recorded and monitored in order to measure progress. The maximum score is 15.

Various terms are used to define altered states of consciousness, as follows.

STUPOR
The patient, although not unconscious and still rousable, exhibits little or no spontaneous activity. The patient may seem to be asleep but will not respond to vigorous stimulation, and will show relatively limited motor abilities, tending to lapse back into sleep when the stimulus ceases. The differential diagnosis is from catatonic schizophrenia and severe depression; the finding of catatonia, posturing of the limbs or flexibilitas cerea is more common with psychiatric disturbances. In organic stupor, the electroencephalograph (EEG) is invariably diffusely abnormal, whereas in psychiatric disease, it will usually be normal.

COMA
This is a condition of absolute unconsciousness judged by the absence of any psychologically understandable response to external stimulus or internal need. It may more simply be defined as a state of ‘unrousable unconsciousness’. The patient will appear to be asleep but is incapable of sensing or responding normally to external stimuli. The condition may vary in depth from the deepest levels, in which there will be no eye-opening, motor or verbal response, to a level in which there may be eye-opening to pain, a weak flexor response of the limbs to pain, and groaning, without recognizable words, in response to pain.

States of altered consciousness will be seen in patients who have suffered from head injury, intoxication with drugs or alcohol, metabolic disturbances or infections or vascular disturbances, and as the result of hypoxic or ischaemic injuries.

Rarely, patients without evident organic illness may appear to have disturbances of consciousness, or pseudocoma. The distinction between pseudocoma and organic disease may be established by testing the oculovestibular reflex in response to the instillation of ice water into the external auditory meatus. In patients in an organic coma, the normal reaction of nystagmus will not been seen, but in patients with pseudocoma, nystagmus will occur, and the patients will usually reveal their responsiveness.

### CONSTIPATION

**ACUTE CONSTIPATION**
Acute constipation may be:
- Due to acute intestinal obstruction
- A symptom of some general disease or of some other acute abdominal disease
- Due to a sudden alteration in daily habits, for example admission to hospital

**Acute intestinal obstruction**
The following points help in the distinction between acute intestinal obstruction and severe cases of acute constipation of other origin:
- In other conditions, the constipation is incomplete, in that flatus – and even a small quantity of faeces – may be passed spontaneously. A rectal examination should always be made. In organic intestinal obstruction, the rectum is usually empty. If it contains faeces, these may be present below an obstruction or, if impacted, may themselves be responsible for the occlusion, but it is exceedingly rare for faecal impaction to produce symptoms quite comparable in severity with those due to acute obstruction. In doubtful cases, it used to be the custom to carry out the two-enema test; the first enema generally brought away a certain amount of faeces, even if obstruction was complete; the second enema, given at an interval of 30 minutes to 1 hour later, resulted in the passage of faeces or flatus if obstruction was incomplete, whereas in complete obstruction the second enema was either retained or expelled unaltered. This test should never be employed; it is exhausting to the patient and time-wasting, and the information obtained is often equivocal. Diagnosis can usually be made on clinical grounds supplemented by abdominal radiographs.
Vomiting is rarely a feature of constipation, whereas it is frequently present in small-bowel obstruction, and in late cases becomes faeculent.

Visible peristalsis, accompanied by noisy borborygmi, is never present except in obstruction.

Obstruction is accompanied by progressive distension of the abdomen.

Pain is usually the first symptom of intestinal obstruction and is colicky in nature; its severity is out of all proportion to the mild abdominal discomfort that may accompany simple constipation.

Plain radiographs of the abdomen are essential in the diagnosis of intestinal obstruction and in attempting to localize its site. A loop or loops of distended bowel are usually seen, together with multiple fluid levels. Small bowel is suggested by a ladder pattern of distended loops, by their central position, and by striations that pass completely across the width of the distended loop and are produced by its circular mucosal folds (Fig. C.17). Distended large bowel tends to lie peripherally and to show the corrugations produced by the taenia coli (Fig. C.18). A small percentage – perhaps 5 per cent of intestinal obstructions – show no abnormality on plain radiographs. This is due to the small bowel being completely distended with fluid in a closed loop, and thus without the fluid levels that are produced by co-existent gas.

Aetiology of acute intestinal obstruction

The causes of intestinal obstruction may be classified as:

- In the lumen:
  - Faecal impaction
  - Food bolus
  - Gallstone ileus
  - Meconium ileus in the neonate
- In the wall:
  - Congenital atresia of the small intestine
  - Crohn’s disease
  - Tuberculous stricture
  - Diverticular disease of the colon
  - Tumours – especially carcinoma of the colon
- Outside the wall:
  - Adhesions and bands (after abdominal surgery or intra-abdominal sepsis)
  - Strangulated hernia (external or internal)
  - Intussusception
  - Volvulus (of the small bowel, or caecal or sigmoid colon)

Before considering any other possibility, all the hernial apertures should be examined, even in the absence of local pain, as a small strangulated femoral hernia in an obese woman, for example, may easily be overlooked.
The following points should be considered in determining the cause of the acute intestinal obstruction.

**Age**  Intestinal obstruction in the newborn should always be suspected in the presence of bile-vomiting: the rectum should be examined first for the presence of an imperforate anus. Other possibilities are congenital atresia or stenosis of the intestine, volvulus neonatorum, meconium ileus and Hirschsprung’s disease. In infants, the most common cause of intestinal obstruction is intussusception, but Hirschsprung’s disease, strangulated inguinal hernia and obstruction due to a band from the tip of a Meckel’s diverticulum should also be considered. In young adults and patients of middle age, adhesions and bands from previous surgery or intraperitoneal inflammation are common, but strangulated hernia and Crohn’s disease are also encountered. In older patients, strangulated hernias, carcinoma of the bowel and diverticular disease, as well as postoperative adhesions, are all common conditions.

**History**  The history of a previous abdominal operation, or of inflammatory pelvic disease in females, strongly suggests the possibility of bands or adhesions. A history of biliary colic or of the symptoms, which may result in cholecystitis, may suggest that obstruction might be due to impaction of a gallstone in the ileum. Obstruction following a period of increasing constipation, perhaps with blood or slime in the stools or spurious diarrhoea, in a middle-aged or elderly patient, suggests cancer or diverticular disease of the colon. The history in an infant or child that blood or mucus have been passed per rectum is suggestive of an intussusception.

**Abdominal examination**  The importance of searching specifically for a strangulated hernia has already been mentioned. The presence of a recent or old laparotomy immediately suggests the diagnosis of postoperative adhesions. Gross distension generally means that the obstruction is in the colon; if occurring very soon after the onset of symptoms, it suggests volvulus of the sigmoid or, less commonly, the caecum. If distension has been present to a lesser extent for some time before the onset of acute symptoms, a growth is likely. In infants and small children, great distension suggests Hirschsprung’s disease. Slight distension occurs when the obstruction is in the duodenum or high in the jejunum.

The diagnosis of intussusception can be made with certainty only when the characteristic sausage-shaped tumour situated somewhere in the course of the colon is felt. In acute obstruction due to cancer, the tumour is often not palpable as it may be disguised by the dilated intestine; however, large masses are sometimes felt, especially when present in the right or left iliac fossa. On the right side, they are generally due to cancer of the caecum, on the left to cancer of the sigmoid colon or diverticular disease.

**Rectal examination**  A growth of the rectum should be recognized easily, although this is rather unusual as a cause of obstruction. Sometimes, a growth of the pelvic colon can be felt through the front wall of the rectum. In infants, the tip of an intussusception may be felt in the lumen of the rectum, and the typical redcurrant-jelly stool (a mixture of blood and mucus) will be seen on the examining finger. Occasionally, the mother will report that a sausage-like structure actually prolapses from the child’s anal verge during the attacks of colic accompanying the intussusception. The present author has only seen this on one occasion. A much-ballooned rectum suggests obstruction in the colon; this is an undoubted fact, but its cause is obscure.

**Vomiting**  The more frequent the vomiting and the earlier the onset of faeculent vomiting, the higher in the intestine is the obstruction likely to be. Its onset is later and its occurrence less frequent in cases of colonic obstruction.

**Symptomatic**

**In acute general diseases**  Constipation beginning acutely is a frequent symptom of a large variety of acute infective and other diseases. It is never so severe as to become a presenting symptom, and the other features in the majority of cases are so much more striking that the presence of constipation has little influence on making a diagnosis.

**In acute abdominal conditions**  Constipation is a conspicuous symptom in most acute abdominal conditions. However, once again, other symptoms are often so well marked that the question of intestinal obstruction hardly arises. Thus, it frequently accompanies acute appendicitis, salpingitis, perforation of a peptic ulcer, and biliary and renal colic. In lead colic, the constipation is not absolute and the occupation of the patient, the blue line on the gums and the presence of punctate basophilia point to the diagnosis.

**Changes in daily routine**  These may precipitate constipation, as in patients admitted to hospital, children going to boarding school or patients suddenly being confined to bed from illness.

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**CONSTITUTION**

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**A**

**B**

**C**

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CHRONIC CONSTIPATION

Constipation can be defined as delay in the passage of faeces through the large bowel and is frequently associated with difficulty in defecation. Most people empty the bowel once in every 24 hours, but there is a considerable range of variation in perfectly normal individuals. In one study of a large working population, this varied from three bowel actions daily to one act every 3 days.

The abnormal action of the bowel in constipation may manifest itself in three different ways:

- Defecation may occur with insufficient frequency.
- The stools may be insufficient in quantity and a certain amount of faeces is retained, although the bowels may be opened once daily or more often (cumulative constipation).
- The bowels may be open daily, yet the faeces are hard and dry owing to prolonged retention in the bowel, dehydration or insufficient residue in the food consumed.

The more common causes of chronic constipation are as follows:

- Organic obstructions, for example carcinoma of the colon or diverticular disease.
- Painful anal conditions, for example fissure-in-ano or prolapsed piles.
- Adynamic bowel, as may occur in Hirschsprung’s disease, senility, spinal cord injuries and diseases, Parkinson’s disease and myxoedema.
- Drugs that decrease the peristaltic activity of the bowel, including codeine, probanthine and other ganglion-blocking agents, and morphine.
- Habit and diet, for example dehydration, starvation, lack of suitable bulk in the diet, and dyschezia.

It is comparatively rare for a patient to consult a doctor on account of constipation without having already attempted to cure themselves with aperients. The symptoms generally ascribed to ‘autointoxication’ caused by intestinal stasis are usually really caused by the purgatives themselves, which may produce depletion of sodium and potassium in the resultant watery stools, or from the abdominal colic and flatulence produced by powerful aperients.

In spite of probable protests, the patient is instructed to see what happens if no drugs are taken for a few days, an attempt being made to open the bowels each morning on a normal diet containing plenty of fruit and vegetables. In most cases, the abdominal pains and so-called ‘toxic’ symptoms disappear. During this test, the bowels are often opened daily, in which case a diagnosis of functional pseudoconstipation can be made, the patient having come to believe, as a result of faulty education combined with advice of friends and with the reading of pernicious advertisements, that he or she was constipated and required aperients to keep well. Frequently, a little psychotherapy in the form of an explanation of the physiology of the bowels and the origin of the symptoms, and persuading the patient to try to open the bowels each morning without artificial help, results in a cure.

The investigation of constipation entails a careful and accurate history and full examination, including an examination of the rectum and sigmoidoscopy, followed, in some cases, by special laboratory tests, a barium enema X-ray examination and colonoscopy.

**Organic obstructions**

The two common causes of narrowing of the lumen of the large bowel are diverticular disease and carcinoma of the colon. Other non-malignant strictures are rare, but they include Crohn’s disease of the large bowel, stricture complicating ulcerative colitis and tuberculous stricture.

**Organic stricture of the colon** is most commonly due to carcinoma. The possibility of cancer should always be considered when an individual above the age of 40, whose bowels have previously been regular, develops constipation of increasing severity without a change of diet or habit, or when a patient who is habitually constipated becomes more so without obvious reason. The constipation is at first intermittent, and may alternate with diarrhoea, or rather with a frequent desire to go to stool without effective evacuation. Aperients become steadily less helpful. There may be colicky pain and episodes of distension, and the patient may notice blood, pus and mucus in the faeces. An examination of the abdomen may reveal a palpable mass due to the presence of the tumour itself, or to inspissated faeces which have become impacted above a cancerous stricture that is itself impalpable. Progressive loss of weight and strength, anorexia and anaemia are rather late features of the disease. A rectal examination reveals usually an empty rectum, but not infrequently a carcinoma in the sigmoid colon can be felt through the rectal wall as the mass in this loop of bowel prolapses into the pelvis. An occult blood test on any faecal material is often positive. Sigmoidoscopy or colonoscopy may visualize the tumour, and biopsy and histological examination can confirm its nature. A barium enema examination is invaluable (Fig. C.19). Diverticular disease of the sigmoid colon can exactly mimic carcinoma, and indeed the surgeon – even at laparotomy – may not be able to differentiate between...
the two conditions. The barium enema examination (Fig. C.20) is often helpful, but the radiologist may have difficulty in distinguishing a stricture due to one or other cause; indeed, these two common diseases not infrequently co-exist. Again, colonoscopy will often be useful in making the differential diagnosis.

Occasionally, extracolonic masses may press upon the rectum or sigmoid colon with resultant constipation, for example the pregnant uterus, a mass of fibroids, a large ovarian cyst or other pelvic tumours.

Painful anal conditions

When defecation is painful, reflex spasm of the anal sphincter may be produced with resultant constipation. A local cause of the pain such as a fissure-in-ano, (see Fig. A.26), strangulated haemorrhoids or a perianal abscess is obvious on careful local examination of the anal verge and surrounds.

Adynamic bowel

In Hirschsprung’s disease (aganglionic congenital mega-colon), there is always a history of constipation dating from the first few months of life. The abdomen becomes greatly enlarged soon after birth, and the outline of distended colon can be seen, often with visible peristalsis. The abdomen finally becomes enormous, and it is then tense and tympanitic (Fig. C.21). There may be eversion of the umbilicus and marked widening of the subcostal angle. The condition is due to the absence of ganglion cells in the wall of the rectosigmoid region of the large bowel, although in some cases a more extensive part of the colon may be involved. Some 80 per cent of patients are male.

A rectal examination reveals a narrow empty rectum above which faecal impaction may be felt; this examination is usually followed by a gush of flatus and faeces. A barium enema examination reveals gross dilatation of the colon leading down to a narrow funnel in the aganglionic rectum (Fig. C.22). Rectal wall biopsy shows complete absence of ganglion cells.

The differential diagnosis is acquired megacolon, a condition of severe chronic constipation, usually commencing at the age of 1 or 2 years, often in a child with mental deficiency. Rectal examination in such cases is typical, impacted faeces being present right up to the anal verge. If necessary, a rectal biopsy may be indicated and will show the presence of normal ganglion cells.

Deficient motor activity of the bowel may be due to senile changes in the elderly, and may be a prominent
feature of myxoedemic patients. Constipation may occur in the course of organic nervous diseases, including Parkinson’s disease, tabes dorsalis, spinal compression from tumour, transverse myelitis and disseminated sclerosis, as well as cord transection in trauma. This is due to disturbance of the motor and sensory pathways responsible for defecation.

Drugs
Many commonly employed drugs have a constipating effect on the bowel; these include codeine, morphine and the ganglion-blocking agents. Constipation accompanied by abdominal pain may be a feature of lead poisoning.

Habit and diet
By far the greatest number of patients complaining of constipation falls into this group. When the faeces are abnormally hard as a result of dehydration, inadequate liquid intake or inadequate cellulose-type material in the diet, rectal examination will reveal impacted faeces of rock-like consistency. This may occur as an acute phenomenon following a barium meal examination, when masses of inspissated barium may lodge in the rectum.

Dyschezia
Dyschezia is the term applied to difficulty in defecation due to faulty bowel habit. The patient ignores the normal call to stool, and the rectum distends with faeces, with the eventual loss of the defecation reflex. The very same patient who gets into this habit is probably one who lives on the modern synthetic diet that is grossly deficient in roughage. As mentioned above, the so-called symptoms of constipation usually result from the purgatives that the patient ingests after becoming anxious about the scarcity of bowel actions. A rectal examination in such individuals often reveals large amounts of faeces in the rectum, and more scybala may be palpated in the sigmoid colon. Dyschezia is, of course, present in those patients who have to remove faeces from the rectum digitally.

Figure C.21 Hirschsprung’s disease. Note the distended abdomen and lack of development. This boy is 15 years old but looks much younger.

Figure C.22 Barium enema in a child with Hirschsprung’s disease.
of corneal transparency, which may be secondary to corneal oedema, cellular infiltration or scarring. Haloes are due to epithelial oedema producing diffraction of the light.

The cornea is examined by ophthalmologists using the slit lamp (biomicroscope) but may be examined with the naked eye by the non-ophthalmologist using focal illumination and magnification (pen-torch and magnifier). Corneal epithelial defects (such as ulcers or abrasions) can be most easily detected by instilling sodium fluorescein, which will stain any epithelial defects. Corneal disease can conveniently be classified into ulcers, which may be infective or immune, nutritional and metabolic disorders, and corneal degenerations.

CORNEAL DEGENERATIONS

The most common degenerative condition of the cornea is arcus senilis, a concentric grey opaque crescent of lipid deposition in the peripheral cornea that, with time, completely encircles the cornea. The ring is about 1–1.5 mm wide and follows the contour of the limbus, but it is separated from it by a clear zone. The outer edge of the ring is a sharp line, but the inner edge shades off gradually. Arcus senilis has no clinical significance except, perhaps, in younger patients in whom it may be associated with serum lipid abnormalities.

Band keratopathy (Fig. C.23) is the name given to characteristic hyaline degenerative changes with calcareous deposits that appear across the interpalpebral area of the cornea in certain eyes. It is rarely found in otherwise healthy eyes, but occurs frequently in blind, shrunken eyes and following prolonged uveitis. Keratoconus (Fig. C.24) or conical cornea is due to an abnormal thinning of the central cornea, and results in a cone-shaped cornea. This condition usually commences around puberty and progresses slowly, although sometimes it develops an acute progressive phase causing sudden visual deterioration and eye discomfort.

CORNEAL ULCERS

The cornea can be infected by viruses, bacteria or fungi. The herpes simplex virus is a common pathogen, producing the typical dendritic ulcer (Fig. C.25). Early lesions comprise opaque epithelial cells that later desquamate to form the branching pattern of the dendritic ulcer which can be demonstrated by fluorescein staining. It is a very painful condition, with much lacrimation and photophobia, and in the absence of proper treatment it tends to last for weeks or even months. It can produce dense stromal scarring that, if centrally situated, leads to visual impairment. The virus can remain dormant but may easily be reactivated in the future by a variety of trigger factors such as poor
general health or exposure to excessive sunlight or topical steroids. Hence, herpes simplex keratitis is frequently a recurrent condition.

The herpes zoster virus, when involving the ophthalmic nerve, can also produce corneal lesions in the form of microdendritic ulcers and corneal opacities. Both herpes simplex and herpes zoster infection can permanently reduce corneal sensation. Corneal sensation can be assessed by touching the cornea with a wisp of cotton-wool and comparing the blink reflex of the two eyes. Damage to the corneal sensory nerves can result in the ulceration of neurotrophic keratitis. The corneal epithelium requires an intact sensory nerve supply to permit normal healing, and neurotrophic ulcers show little tendency to heal, often requiring a tarsorrhaphy.

Bacterial ulcers or abscesses often follow minor trauma to the cornea. The most common pathogens are *Streptococcus pneumoniae* and *Staphylococcus aureus*, but *Pseudomonas*, *Proteus*, *Klebsiella*, *Escherichia coli* and *Neisseria* can also be causative. Contact lens wear has become a major risk factor in the causation of corneal ulcers and infectious keratitis. This is particularly so when there is poor contact lens care and poor hygiene, especially with extended-wear contact lenses. Bacteria cultured from contact lens-associated corneal infections are commonly *Pseudomonas aeruginosa* and *Staphylococcus aureus*. Clinically, the bacterial ulcers appear as yellowish irregular opacities associated with corneal stromal necrosis. They are usually associated with reactive iritis with an outpouring of white blood cells into the anterior chamber; these cells settle inferiorly to form a pus level or hypopyon (Fig. C.26). The causative organism is identified and the appropriate antibiotics given. The ulcer may heal without perforating, but if the resulting scar is both large and centrally placed, the vision is ultimately often much impaired. If perforation does occur, the underlying iris may adhere to the site of perforation with subsequent dense corneal scar formation, which may then distend to form a bulging anterior staphyloma.

*Marginal ulcers* situated around the periphery of the cornea, with a clear zone between the ulcer and the limbus, tend to remain superficial. They most commonly follow *Staphylococcus aureus* conjunctivitis or blepharitis, and are produced as an immunological reaction against staphylococcal exotoxins. *Interstitial keratitis* is a late manifestation of congenital syphilis that produces stromal corneal opacities with the associated stigmata.

Keratitis due to *Acanthamoeba*, a genus of amoeba found in fresh water and soil, is a serious infection of the cornea that is occurring more commonly in contact lens wearers who use fresh water as opposed to proper contact lens solutions. Acanthamoebic infections can masquerade as herpetic or fungal keratitis in particular, and the pain is often out of proportion to the clinical signs. *Keratomalacia* due to vitamin A deficiency is characterized by desiccation and subsequent necrosis of the cornea and conjunctiva. It affects undernourished infants and is extremely rare in the UK. Urgent administration of vitamin A is required, together with attention to the corneal hydration.

**COUGH**

Alex West

Healthy persons seldom cough; their scant bronchial secretions result in a thin sheet of mucus that is constantly carried up the tracheobronchial tree towards the larynx by the action of cilia. On reaching the pharynx, the secretions raised in this way are disposed of into the alimentary tract by unconscious acts of swallowing.

Coughing is an essential defence mechanism that protects the airways from the adverse effects of inhaled noxious substances and also serves to clear them of retained secretions. Patients recognize that coughing indicates an abnormality, and this symptom is one of the most frequent reasons given for seeking medical advice. Coughing may be produced voluntarily, but more often it results from reflex stimulation. To a lesser extent, it can be suppressed voluntarily. The involuntary initiation of a cough takes place in a reflex arc. Extrathoracic cough receptors are present in the nose, oropharynx, larynx and upper trachea. Intrathoracic
irritant receptors are located in the epithelium of the lower trachea and large central bronchi, which are the air passages from which coughing effectively expels retained secretions or foreign material. Efferent pathways include the recurrent laryngeal nerves to cause closure of the glottis, and the corticospinal tract and peripheral nerves to cause contraction of the thoracic and abdominal musculature. The cough receptors may accommodate to repeated stimuli – as they often do in cigarette smokers, who may only cough after the first cigarette of the day. The cough reflex becomes less sensitive in the elderly, and it is lost in anaesthesia and unconsciousness, leading to an increased danger of aspiration pneumonia.

The act of coughing occurs in three phases. The first phase is a preliminary deep inspiration, and the second is closure of the glottis, relaxation of the diaphragm and contraction of the thoracic and abdominal expiratory muscles, generating a positive pressure of 100–300 mmHg within the thorax. Because the positive pressure in the pleural space is higher than the luminal pressure in the trachea and central bronchi, a pressure difference is created that causes the posterior membranous portion of the airway walls to fold inwards and partially obliterate the lumen. When the third phase occurs – namely, sudden relaxation of the glottis – the linear velocity of airflow through the narrow channels is markedly increased, creating forces that dislodge secretions and particles from the mucosal surface. During cough, the volume rate of flow out of the lungs (litres per second) is very similar to that obtained during a forced expiratory manoeuvre – a fact that is not always appreciated. In patients with severe airflow obstruction, high rates of flow cannot be generated because the airways are already narrowed; such patients may have prolonged wheezy coughs that sometimes cause the involuntary effects of a Valsalva manoeuvre, and resultant cough syncope, which is occasionally accompanied by convulsions mimicking epilepsy.

In general, the diagnosis of the cause of cough depends not only on an analysis of the cough itself but also on the other symptoms and physical signs and, above all, the chest radiograph. When the chest radiograph shows a significant abnormality such as lobar collapse (Figs C.27–C.29), carcinoma (Fig. C.30), bronchopneumonia (Fig. C.31) or pulmonary tuberculosis (Figs C.32 and C.33), the reason for the cough is established and the next step is to initiate appropriate treatment. Diagnostic problems arise in those patients in whom the chest radiograph appears normal. The disorders that need to be considered are listed in Table C.3. Some helpful diagnostic clinical clues and signs and further investigations are listed in the same table. Enquiry about the presence or absence of sputum, whether the cough is occasional or...
persistent, or is provoked by any activity or situation, and especially whether there is disturbed sleep, will provide helpful pointers to the diagnosis. When the causes of the chronic cough are analysed, asthma frequently heads the list, and indeed chronic non-productive cough—especially at night—may be the sole presenting complaint of patients subsequently proven to have bronchial asthma. There are two other main causes of chronic ‘idiopathic’ cough in non-smokers with normal chest X-rays: acid reflux, which can be a cause even if symptoms of heartburn are minimal, and this classically does not occur at

Figure C.27 Chest radiograph showing a left upper lobe collapse due to a bronchial carcinoma. A veil-like opacity is obscuring the left heart border (white arrow).

Figure C.28 Chest radiograph showing right upper lobe collapse in a child (white arrow).

Figure C.29 Chest radiograph showing right upper lobe pneumonia due to tuberculosis (white arrow), with enlarged lymph nodes (black arrows).

Figure C.30 Chest radiograph (a) and computed tomography scan (b) showing a lung cancer at the left hilum invading the left pulmonary artery (white arrow).
night; and posterior nasal drip. These three can be remembered as the 3Rs – Reactive airways (asthma), Reflux and Rhinitis.

Dry cough is also a feature of lung fibrosis from any cause. Typically, the chest radiograph will be abnormal but, if equivocal, thoracic high-resolution computed tomography scanning is helpful in confirming or refuting the presence of structural lung disease.

An intractable dry cough is now recognized as an important side effect of angiotensin-converting enzyme inhibitors.

In pertussis, the characteristic cough is paroxysmal and occurs in bouts that may last for a minute or two and culminate in vomiting, in addition to a characteristic terminal inspiratory whoop. In severe paroxysms, the child may become cyanosed. On examination, the most striking finding in the chest is a negative one, the rhonchi characteristic of ordinary acute bronchitis being generally absent. A sublingual ulcer on the frenum linguae due to the friction of the protruding tongue on the lower front teeth during long paroxysms of coughing is a helpful finding, as is a history of exposure to infection.

A cough that an adult patient has had intermittently since childhood is more likely to be associated with bronchiectasis than one for which the onset can be dated more recently.

In children and even young adults with a persistent or recurrent cough, the possibility of cystic fibrosis must be borne in mind; the presence of associated bowel symptoms should lead to an estimate of sweat sodium levels.

A cough appearing and persisting for the first time in a young adult – especially if of Asian origin – must give rise to suspicion of pulmonary tuberculosis and calls for further investigation.
for complete investigation including a chest radiograph and sputum examination. Similarly, a cough appearing for the first time in middle age – especially in a man – must raise suspicion of bronchial carcinoma, which should only be dismissed after a complete investigation.

The characteristic morning cough of the cigarette smoker is due to a chronic pharyngotracheobronchitis; many cigarette smokers regard it as a normal part of their lives, and refer to it as ‘clearing the throat’. It is in fact the first symptom of chronic bronchitis. The frequency with which chronic bronchitis and bronchogenic carcinoma co-exist – both being directly associated with tobacco consumption – has led to the important axiom that any change in the character or pattern of a chronic cough warrants investigation for carcinoma of the lung. Chronic nasal sinusitis may produce or contribute to this symptom, secretions that have trickled down into the trachea during sleep being expelled when the patient wakes. A cough that appears on first lying down at night or on some other changes of posture is suggestive of localized bronchiectasis or chronic pulmonary suppuration. In older people, recurrent aspiration is a not uncommon cause that is often overlooked. It may be related to oesophageal regurgitation, stricture or neurological disease affecting swallowing. Asthma and cardiac failure also commonly cause cough in the elderly, but this may be especially prominent at night. Asthmatic patients sometimes complain of cough as a predominant feature of their attacks, and wheezing and dyspnoea may be relatively trivial. In typical asthma, cough may also occur on exercise or nocturnally, waking patients from sleep. A cough with dyspnoea or orthopnoea waking the patient from sleep may also be due to pulmonary congestion or oedema due to left ventricular failure in hypertension, aortic valvular disease or disorders of the myocardium. Because such nocturnal attacks of paroxysmal dyspnoea in these conditions may be accompanied by wheeziness, they are sometimes referred to as ‘cardiac asthma’. The important findings that might help to differentiate bronchial and cardiac asthma are listed in Table C.4.

The presence or absence of expectoration and the quality of expectoration are important diagnostic features of any cough. It should be remembered that many patients find it difficult to expectorate, and they habitually expel small amounts of secretion through the larynx by cough and then swallow it. This is the rule in children. A dry cough, or one producing only scant mucoid sputum, may be due to inflammation or tumour of the larynx, when there will be associated hoarseness. Laryngeal involvement in syphilis is a very unusual cause of a dry cough with hoarse voice. Another cause of cough associated with a weak voice often described as ‘hoarseness’ is carcinoma of the bronchus with recurrent laryngeal nerve involvement. A dry cough may be a manifestation of nervousness, but it should not be accepted as such without proper investigations. A dry cough may also be due to external pressure on a bronchus by a mediastinal mass such as benign or malignant tumour, or by enlarged mediastinal lymph nodes due to reticulosis or tuberculosis; the latter should be particularly considered in those of Asian origin. Cough due to external pressure on the trachea is usually described as ‘brassy’ or ‘bovine’.

If a cough is productive, the quality and mode of production of the sputum should be noted. Frankly purulent sputum suggests bronchiectasis, lung abscess, being primary or secondary to bronchial obstruction by new growth, foreign body or cavitating pulmonary tuberculosis. The odour of the sputum is important: a malodorous sputum almost invariably indicates infection with anaerobic organisms and suggests bronchiectasis, inhalation pneumonia, lung abscess or bronchopleural fistula with expectoration of a putrid empyema (see also SPUTUM, p. 636). Cough due to a bronchopleural fistula is characteristically

### Table C.4 Differentiation of bronchial and cardiac asthma

<table>
<thead>
<tr>
<th>Bronchial asthma</th>
<th>Cardiac asthma</th>
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<tbody>
<tr>
<td>History of previous bronchial asthma</td>
<td>History of hypertension, valvular or ischaemic heart disease</td>
</tr>
<tr>
<td>Often young – sometimes old</td>
<td>Rarely young – usually old</td>
</tr>
<tr>
<td>Expiratory wheezing</td>
<td>Sometimes also expectoratory wheezing</td>
</tr>
<tr>
<td>No crackles</td>
<td>Many basal crackles</td>
</tr>
<tr>
<td>ECG normal</td>
<td>Gallop rhythm / cardiac failure</td>
</tr>
<tr>
<td>Chest radiograph shows hyperinflation of lungs</td>
<td>Evidence of pre-existing cardiac disorder</td>
</tr>
<tr>
<td>Bronchial hyperactivity (peak expiratory flow rate measurements)</td>
<td>No evidence of bronchial hyperactivity</td>
</tr>
<tr>
<td>Methacholine or histamine challenge</td>
<td></td>
</tr>
</tbody>
</table>
dependent on position; the cough is worse when the patient lies on their good side and is relieved by lying on the side of the lesion.

Coughing seems to provoke more coughing. Paroxysms of coughing as in pertussis may terminate in vomiting, which seems to break the cycle. Paroxysmal attacks may also terminate in syncope. At times, severe coughing attacks have continued to the point of utter exhaustion. The muscular force developed during coughing may be sufficient to cause occasional fractures of the ribs (cough fractures) and even compression fractures of the vertebral bodies. In some cases, no physical cause for the cough may be detected, and in some of these patients psychogenic factors are important.

**CRAMPS**

Mark Kinirons

Cramps are involuntary, painful, sudden spasms of voluntary muscles that become hard and ‘knotted’. Cramps may be initiated by a sudden voluntary contraction, but they usually occur at rest. They may persist for seconds or minutes, and can generally be relieved by a slow passive extension of the affected muscle. Electromyography of a muscle in cramp shows fluctuating high-frequency bursts of motor unit activity. Cramps are a common symptom, and only infrequently indicate disease, particularly if they occur in the legs. Underlying medical causes, however, include salt depletion, hypocalcaemia, muscle ischaemia, myopathy and neurological disease.

Cramps may occur in normal people under certain physiological (rather than disease) conditions. They often occur during pregnancy. Excessive unaccustomed exercise can cause cramps; thus, athletes and swimmers commencing training are often affected, the cramps disappearing with increasing fitness. Cramp of the intercostal muscles (stitch) may occur during and after exercise in untrained individuals. Excessive use of combinations of small muscle groups can cause cramps (e.g. writer’s or musician’s cramp). These are focal dystonias (without generalized neurological disease) that make it hard to perform that particular motor task. Medications and psychotherapy are generally unhelpful, but botulinum toxin has been shown to be effective (its use should, however, be justified by the severity of the condition). Night cramps typically occur in older people and may be painful enough to wake patients from sleep. The foot and calf muscles are most commonly affected, the thighs less often. Quinine is moderately effective at preventing nocturnal leg cramps.

Chronic muscle ischaemia due to atherosclerosis causes intermittent claudication, most commonly in the calves, where walking a certain distance (often consistent in any one individual) causes a cramping pain forcing the person to stop. Rest will tend to relieve the pain, and patients with claudication are advised to increase their daily exercise to at least four half-hour walks each week. Signs of disease are absent lower-limb pulses, dusquines of the feet on hanging the legs down, atrophic skin and, if severe, cold white leg and punched-out ulcers.

Hypocalcaemia causes cramp, most strikingly seen in the forearm and hand muscles as carpopedal spasm (tetany). Hypoparathyroidism, malabsorption and inadequate dietary calcium intake can cause hypocalcaemia. Acute alkalosis from hysterical hyperventilation can also lower ionized calcium in plasma and cause carpopedal spasm. Chronic renal failure may also result in hypocalcaemia and, with hypomagnesaemia, be a cause of cramp. In these patients, cramps may be particularly troublesome during dialysis. Salt depletion as a result of hard physical work in a hot environment, profuse sweating and hypertonic fluid replacement (i.e. water) causes heat cramps, typically presenting as severe spasms of the extremities. Rectal temperature is usually normal, and treatment consists of rest and salt replacement (preferably by food and fluids containing sodium chloride, rather than by salt tablets). Cramps may also occur from salt and water imbalance due to diarrhoea and vomiting, burns, diuretic use, fistula and small-bowel obstruction.

Neurological diseases affecting the spinal cord or peripheral nerves can cause cramps. Flexor spasms of the legs may be seen in patients with spinal cord injury and myelitis due to multiple sclerosis. Diseases of the peripheral nerves such as alcoholic or diabetic neuropathy tend to cause cramps of the legs and feet, whereas forearm cramps raise the possible diagnosis of motor neurone disease.

In addition to diuretics, drugs associated with cramps are beta-adrenoceptor stimulants (e.g. terbutaline), steroids, calcium-channel antagonists (e.g. nifedipine), statins (myopathy) and phenothiazine-type neuroleptics. The latter cause a particular form of cramp and spasm of the facial and neck muscles (orofacial and nuchal dyskinesia).

Very rarely, cramps may be due to enzyme disorders of muscles, for example phosphoglycerate mutase, phospho-glycerate kinase and adenylate deaminase deficiency. Tetanus is a form of cramp due to infection.
by *Clostridium tetani*. The spasm appears first in the facial muscles and then spreads over the body. Cramps due to *lead poisoning* or *strychnine* are rarely seen and are only of historical interest.

**Crepitus**

Mark Kinirons

Crepitus is a term generally used to denote the grating or crackling sensation and noise produced when two rough substances rub together, for instance the grating that can be felt and heard between the fractured ends of a bone. Crepitus from within a joint should be distinguished from sharp cracking sounds occurring in normal or abnormal joints after temporary immobility, and from the slipping of tendons and ligaments over bone surfaces on movement.

Articular crepitus arises most commonly from osteoarthrotic degenerative changes in joints, most commonly the knee, where subpatellar fine friction can be felt with the hands in many cases, often early in the degenerative process. The patient should be reassured that it is not a serious finding and is felt by most middle-aged or elderly persons at some time in their lives.

A common and unimportant observation after middle age is the awareness of a grating sensation in the neck, apparent to both patient and clinician. In acromegaly, gross crepitus throughout the full range of articular movement is often found in large peripheral joints, as it may be in other syndromes associated with hypermobile joints, such as Marfan’s and Ehlers–Danlos syndromes. A coarser crepitation may be felt after trauma and in chondromalacia patellae. Coarse crepitation or ‘crackling’ may be felt with the hands or heard with the ear wherever degenerative changes occur, for instance in the neck or shoulder, but is more common in the more mobile weight-bearing joints, where cartilage has become worn and degenerated, as in, most commonly, the knee.

In rheumatoid arthritis, a fine ‘rubbing’ crepitus occurs, which can be felt over the joints in relatively early cases; this may become coarser as the disease advances. At the shoulder, inflammatory or degenerative processes may produce crepitus at the scapulohumeral or acromio-clavicular joints, and a peculiar sensation may be experienced with the hand over the scapular border as though the bone were moving over soft marbles; this, again, is not a sign of serious disease, and the patient can, in most cases, be fully reassured.

On palpation of a joint with hydrarthrosis, the so-called ‘silken crepitus’ can occasionally be felt, as if two silken surfaces were being rubbed together by the examiner’s hands. *Tenosynovitis*, especially around the flexor tendons at the wrist, can also produce a feeling of crepitus to both patient and examiner. When there is an enlargement of a bone without fracture, and when on palpation a feeling of crepitus or eggshell cracking is obtained, it indicates that some tumour is eroding the overlying cortex. Radiography will be necessary to establish the diagnosis. Rarefraction of the bones of the skull, either as the result of syphilitic lesions in adults, or of *hydrocephalus* or *craniotabes*, especially in the occipital region of congenitally syphilitic and rickety infants, may make the skull bones so thin that they bend readily on pressure, and sometimes the result is a sensation of crepitus. The diagnosis is generally obvious. The condition is very rare.

Apart from bony, arthritic or synovial changes, a characteristic feeling of crepitus may be felt beneath the skin when gas or air has accumulated in the subcutaneous tissues as the result of surgical emphysema.

**Crusts**

Barry Monk

Crusts or scabs are secondary skin lesions formed when serum, blood, sebum, purulent exudate or a mixture of these dries on the surface of the skin (Fig. C.34). In addition, they may contain dirt and the remains of local applications (e.g. calamine lotion). It is important to distinguish crusts from scales (see SCALY ERUPTIONS, p. 599), the causes being quite distinct. Crusts vary considerably in thickness, from the light crust of dermatitis to the thick barnacle-like scabs seen in keratoderma blennorrhagica. It may sometimes be necessary to remove crusts by prolonged soaking in

![Figure C.34 Actinic keratoses.](image-url)
order to make a diagnosis of the underlying lesion. This is particularly important when a crust overlies a neoplasm (e.g. a rodent ulcer on the scalp) (Box C.6). Impetigo is the classic cause of golden-coloured crusts on the skin (Fig. C.35). These are most common around the mouth and face of children, but they can be very widespread or affect any age group. Acute vesicular ‘wet’ eczema results in crust formation as the exudates dries; if this becomes golden-coloured, secondary impetiginization due to infection with Staphylococcus aureus should be suspected (Fig. C.36). Herpes simplex sores become crusted on the fourth or fifth day, and crusting is prolonged if there is a secondary impetigo. The same may occur in Herpes zoster, but here crusts are often haemorrhagic. In eczema herpeticum (Kaposi’s varicelliform eruption), crusting can be very extensive and pose a difficult nursing problem. Pemphigus (Fig. C.37) is the bullous disorder that is characterized by crusting and erosions because the blisters are superficial and therefore very fragile. A search should be made for the characteristic mucosal erosions. These crusts are notoriously slow to heal and are probably the only situation where the application of a potent topical corticosteroid can speed re-epithelialization.

Scabbing can be profuse around stasis ulcers, particularly if a dermatitis medicamentosa (e.g. to lanolin or topical antibiotic) has developed. The crusting over pyoderma gangrenosum overlies the characteristic cribri-form fibrous scarring.

In keratoderma blennorrhagica, the skin eruption of reactive arthritis, crusty rupioid nodules may be seen on the limbs accompanying the pustular crusted lesions on the palms and soles. In the rupioid form of secondary syphilis, the crusts are greenish or blackish and consist of several layers, each smaller than the one immediately below it, so that a pyramidal structure is formed resembling a barnacle. Removal of these crusts

<table>
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<tr>
<th>Box C.6</th>
<th>Crust</th>
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<tr>
<td>Impetigo</td>
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</tr>
<tr>
<td>Eczema</td>
<td></td>
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<tr>
<td>Trauma – thermal, chemical, factitial</td>
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<td>Herpes simplex, eczema herpeticum</td>
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<tr>
<td>Herpes zoster</td>
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<tr>
<td>Pemphigus</td>
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<td>Stasis ulcers</td>
<td></td>
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<tr>
<td>Pyoderma gangrenosum</td>
<td></td>
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<tr>
<td>Keratoderma blennorrhagica (reactive arthritis)</td>
<td></td>
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<tr>
<td>Syphilis – rupioid</td>
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<tr>
<td>Yaws</td>
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Central cyanosis is the hallmark of the ‘blue and previously termed chronic bronchitis and emphysema). chronic obstructive pulmonary disease (COPD, common respiratory cause of central cyanosis is severe, will result in central cyanosis. The most Virtually any diffuse lung disease, if sufficiently usually in association with hypercapnia. hypoventilation will also result in arterial hypoxaemia, blood to the systemic arterial circulation. Alveolar tension and will contribute poorly oxygenated but well-perfused alveoli will have a low oxygen matching of ventilation to perfusion. Poorly ventilated due to respiratory disease is the impairment of Central cyanosis is due either to arterial hypoxaemia or to the presence of the abnormal pigments methaemoglobin or sulphamoglobin. Central cyanosis is seen in the mucous membranes, particularly the tongue, and in good lighting can be identified by most observers when the arterial oxygen saturation falls below about 87 per cent. In anaemia, greater desaturation will occur before cyanosis is apparent, as less reduced haemoglobin will be present at a given saturation. Peripheral cyanosis is due to reduced blood flow in the skin, allowing more oxygen to be extracted than normal. Peripheral cyanosis may result from local vasoconstriction, from a low cardiac output or from a combination of both. It is apparent in exposed areas such as fingers, ears, nose and cheeks, but not in the mucous membranes or the warmer parts of the skin.

**CENTRAL CYANOSIS**

**Respiratory disease**

The most common mechanism for arterial hypoxaemia due to respiratory disease is the impairment of matching of ventilation to perfusion. Poorly ventilated but well-perfused alveoli will have a low oxygen tension and will contribute poorly oxygenated blood to the systemic arterial circulation. Alveolar hypoventilation will also result in arterial hypoxaemia, usually in association with hypercapnia. Virtually any diffuse lung disease, if sufficiently severe, will result in central cyanosis. The most common respiratory cause of central cyanosis is chronic obstructive pulmonary disease (COPD, previously termed chronic bronchitis and emphysema). Central cyanosis is the hallmark of the ‘blue and bloated’ form of COPD, and it is then due to a combination of ventilation–perfusion mismatching and hypoventilation. Patients with the ‘pink and puffing’ form of COPD have impaired ventilation–perfusion matching but avoid marked arterial hypoxaemia by maintaining adequate ventilation at the cost of developing dyspnoea. Central cyanosis is a feature of advanced pulmonary fibrosis from whatever cause when it results from ventilation–perfusion mismatching. Ventilation–perfusion mismatching is also the cause of cyanosis in pulmonary embolism, pulmonary oedema and pneumonia.

Alveolar hypoventilation is the cause of central hypoxaemia following excess sedation and in patients with ventilatory failure due to impaired chest wall movement. Impaired chest movement may result from kyphoscoliosis or from paresis of the respiratory muscles. Alveolar hypoventilation may also cause cyanosis in patients with sleep apnoea/hypopnoea, a condition characterized by daytime sleepiness and loud snoring, or the obesity hypoventilation syndrome, a condition characterized by obesity and episodes of hypoventilation occurring principally at night, leading to type II respiratory failure and consequent hypoxia, the ‘Pickwickian syndrome’ (see SLEEP, DISORDERS OF, p. 621). Alveolar hypoventilation also occasionally occurs following obstruction of a major airway. For example, the larynx may be obstructed by the aspiration of a foreign body, by laryngeal oedema (usually due to angioneurotic oedema) or by bilateral abductor paralysis following thyroidectomy. Tracheal obstruction from thyroid carcinoma, haemorrhage into a thyroid cyst, malignant lymph nodes, primary tracheal tumours and benign tracheal stenosis following tracheostomy or intubation are all rare causes of cyanosis due to alveolar hypoventilation.

**Cardiovascular disease**

Although the most common cardiac cause of mild hypoxaemia is left heart failure, usually on the basis of ischaemic heart disease, this is rarely severe enough to cause cyanosis except during acute attacks of pulmonary oedema. Although cyanotic congenital heart disease is a less common cause of hypoxaemia, it is a much more common and more dramatic cause of cyanosis. An important point differentiating cyanosis due to a central veno-arterial shunt from that due to lung disease is the response to breathing oxygen. In lung disease, cyanosis will disappear and arterial hypoxaemia greatly improve as a result of breathing 100 per cent oxygen, unless many perfused alveoli are completely unventilated. In cyanotic congenital heart disease, oxygen produces little change in arterial
oxygen saturation, as the blood in the shunt does not participate in gas exchange. Other features common to all types of cyanotic congenital heart disease are clubbing of the fingers and toes, and polycythaemia. In severe cases, the haemoglobin concentration may be well over 20 g/dl.

Fallot’s tetralogy is by far the most common congenital cardiac lesion to cause cyanosis. The four classic features are pulmonary stenosis, ventricular septal defect, overriding aorta and right ventricular hypertrophy; of these, the last is the direct haemodynamic consequence of the pulmonary stenosis; the overriding aorta, although characteristic, is not a major factor in determining the presence or severity of cyanosis. An associated anomaly in several cases is a right-sided aortic arch; this must not be confused with dextro-position of the aorta – a synonym for overriding. Cyanosis is not often present at birth but develops, with clubbing, during the first year or two of life. The symptoms include dyspnoea, relieved by squatting, and syncope associated with deepening of the cyanosis (see DYSPNOEA, p. 144; FAINTS, p. 195). The arterial and venous pulses are unremarkable except for, occasionally, a dominant ‘a’ wave in the jugular venous pulse. The apex beat is little displaced, and there may be a slight or moderate right ventricular impulse at the left sternal border. The systolic murmur is rather short, and often loud enough to be accompanied by a thrill; the second sound is single. The definitive diagnosis can be made by echocardiography or angiocardiography.

Pulmonary atresia is embryologically related to Fallot’s tetralogy and usually produces very severe cyanosis. The differentiation is easily made by the absence of the systolic murmur of pulmonary stenosis in atresia and its replacement by a continuous subclavicular murmur, usually bilateral, due to large anastomoses between the bronchial and pulmonary arteries. Other conditions that may be more difficult to distinguish clinically from Fallot’s tetralogy are pulmonary stenosis with reversed interatrial shunt and Eisenmenger’s complex, properly so-called, that is pulmonary hypertension with a reversed interventricular shunt. Tricuspid atresia is another cause of cyanosis, characterized by an unusual electrocardiographic pattern suggesting right atrial and left ventricular hypertrophy.

Transposition of the great arteries is a relatively common form of cyanotic congenital heart disease in infancy, although few untreated patients survive. The aorta arises from the right ventricle and the pulmonary artery from the left and, clearly, communications between the two circulations, at atrial or ventricular level, must also be present if life is to be maintained. Pulmonary stenosis or pulmonary hypertension may also be present to complicate the haemodynamic situation. The physical signs vary with the associated abnormalities, but the X-ray may be of diagnostic value. The vascular pedicle of the heart is rather narrow in the posteroanterior view, and pulmonary plethora is nearly always present. As oligaemia of the lung fields is the rule in most types of cyanotic congenital heart disease, the association of cyanosis with plethora is very suggestive of transposition. Angiocardiography is diagnostic, showing, in the lateral view, the aorta arising anteriorly from the right ventricle, and the pulmonary artery posteriorly from the left. Very rare types of cyanotic congenital heart disease include drainage of the superior or inferior vena cava into the left atrium and some cases of Ebstein’s anomaly of the tricuspid valve with reversed interatrial shunt.

Pulmonary arteriovenous malformations may cause central cyanosis if the shunt is large enough. They are frequently associated with cutaneous telangiectases, especially on the lips, and may occur in the familial condition hereditary haemorrhagic telangiectasia. Continuous murmurs may be heard over the aneurysms, which may be multiple. In the radiograph, one or more round opacities are visible, and a computed tomography scan will often reveal the feeding artery and draining vein.

Pigmentary cyanosis

Cyanosis is a major feature of a group of conditions in which ferric (Fe³⁺) rather than ferrous (Fe²⁺) iron is present in haemoglobin, producing methaemoglobin. Sulphaemoglobin is a poorly characterized substance that can be produced by the action of hydrogen sulphide on haemoglobin. Small amounts of these pigments, for example, 1.5 g of methaemoglobin or 0.5 g of sulphaemo-globin, are said to produce as marked cyanosis as 5 g of deoxygenated haemoglobin. Methaemoglobinemia may be congenital or acquired. Congenital methaemoglobinemia is due to deficiency of NADH-methaemoglobin reductase have a recessive inheritance. Methaemoglobinemia may also result from congenital abnormalities of the alpha-or beta-globin chain of haemoglobin, rendering the molecule unresponsive to methaemoglobin reductase, and these varieties may be inherited as autosomal dominant characteristics. Acquired methaemoglobinemia can result from the ingestion of a large group of chemicals including oxidizing agents such as nitrates and chlorates. Nitrates can be converted to nitrites in the bowel, and cyanosis has been reported in infants in association with a high nitrate content in their
drinking water. Poisoning with potassium chlorate can also produce methaemoglobinaemia, apart from its more serious effects. Aniline dyes have also been incriminated; these compounds can be absorbed through the intact skin. Phenacetin can also cause methaemoglobinaemia and cyanosis, although, due to declining use of this preparation, this is now rare. There has been an increase in the incidence of methaemoglobinaemia related to the use of dapsone in HIV. The symptoms of methaemoglobinaemia are those of severe anaemia, to which the condition is closely analogous, namely dyspnoea on exertion and dizziness. Clinicians should be alerted to the presence of methaemoglobinaemia by an apparent oxygen saturation of 85 per cent on pulse oximetry that does not improve on an increased concentration of inspired oxygen. The diagnosis can be confirmed by co-oximetry, available on most modern blood gas analysers, which will detect the characteristic absorption of methaemoglobinaemia.

Sulphaemoglobinaemia is a poorly defined condition that may occur in conjunction with methaemoglobinaemia following the ingestion of nitrates, nitrites, aniline dyes and phenacetin. It has also been reported in conjunction with chronic constipation and with malabsorption. There is no congenital form of this condition. Sulphaemoglobinaemia is usually asymptomatic.

PERIPHERAL CYANOSIS

When red cell transit through cutaneous vessels is delayed, continued oxygen extraction will decrease the oxygen saturation of the haemoglobin, and cyanosis will appear. This may result either from increased resistance to blood flow, from decreased cardiac output or from increased blood viscosity.

Increased resistance to blood flow

The most common cause of peripheral cyanosis is the transient and appropriate vasoconstriction in response to cold. The best-known medical conditions associated with peripheral cyanosis are Raynaud’s disease and Raynaud’s phenomenon, which are described in detail in the section on FINGERS, DEAD (WHITE, COLD) (p. 201). Acrocyanosis is a relatively benign condition due to spasm of the smaller cutaneous arteries and arterioles. The hands and fingers are cold and mottled red and blue, but pain is not a feature. Arteriolar spasm is also the mechanism of erythrocyanosis, a disease almost confined to young women, in which cyanotic blotches are seen in the lower parts of the legs. Cyanosis of the affected leg or legs occurs occasionally following deep venous thrombosis, particularly if collateral drainage is poor. This condition has been termed phlegmasia caerulea dolens. Similarly, cyanosis of the face may occur, together with gross venous engorgement and oedema, as a feature of superior vena caval obstruction from whatever cause. This mechanism may contribute to the so-called traumatic cyanosis following crush injury to the chest, although hypoxaemia due to lung trauma may contribute.

Decreased cardiac output

The ashen grey appearance of patients with severe shock is a typical example of peripheral cyanosis due to a marked fall in cardiac output, and it occurs irrespective of the cause of the shock. The low cardiac output and vasoconstriction that occur with left ventricular failure can also cause peripheral cyanosis, although central cyanosis will usually co-exist. Similarly, cyanosis in massive pulmonary embolism is usually both central and peripheral. The classical malar flush in mitral stenosis is an example of local peripheral cyanosis, but why this should be so sited is unknown.

Increased blood viscosity

Cyanosis may occur in polycythaemia rubra vera despite a normal arterial oxygen saturation. This is presumed to result from decreased cutaneous blood flow due to increased viscosity.
Deafness is rarely total or complete. The term ‘hearing loss’ is a better term to use as it implies that there may be degrees of deafness. Hearing loss is an extremely common problem, and constitutes one of the major handicaps affecting mankind. Most of us will have experienced a temporary hearing loss in one or other ear at some time, perhaps when flying or associated with a cold, and we should be able to appreciate how incapacitating permanent and more severe loss of hearing must be. In fact, almost 60 per cent of the population will have acquired a significant degree of hearing loss by the time they retire from full-time employment.

From a clinical standpoint, three types of hearing loss are recognized, namely conductive, sensorineural and mixed.

- **Conductive hearing loss** is caused by lesions in the external and/or middle ear that attenuate or prevent sound reaching the cochlea.
- **Sensorineural hearing loss** is caused by lesions within the cochlea or affecting the auditory nerve and/or higher pathways.
- **Mixed hearing loss** is caused by a combination of conductive and sensorineural elements.

**HEARING TESTS**

These tests are undertaken to determine the nature and severity of hearing loss. Both psycho-acoustic and objective tests have been devised and are employed in clinical practice.

**Voice tests**

An initial assessment of hearing loss can be gained by occlusion of the contralateral ear while speaking or whispering into the ipsilateral ear. The patient should be told to repeat exactly what the examiner says and be positioned so that they cannot see the examiner’s lips. Anyone with normal hearing will be able to hear a whispered voice at a distance of 60 cm. Hearing in the contralateral ear is usually masked by gentle pressure on the tragus, or by holding paper over the ear and scratching it when speaking. An alternative and better method of masking the non-test ear is by use of a Barany box; this is a sound-generating device that is inserted into the contralateral ear canal and emits a loud noise while testing the other ear. More accurate measures of the spoken voice can be obtained with sound pressure meters.

**Tuning-fork tests**

With these simple, chair-side tests, it is possible to distinguish between conductive and sensorineural hearing losses. Only 256 cycles per second (cps) or 512 cps tuning forks should be used. Lower-frequency tuning forks are appropriate for vibration sense tests only.

**The Rinne test**

The tuning fork is struck and its base held firmly on the patient’s mastoid process until they no longer hear the tone. The fork is then rapidly transferred so that the vibrating forks are close to the external auditory meatus. If the patient continues to hear the sound, it is considered that they hear better by air conduction than by bone conduction – a positive test result. In fact, in patients with normal middle-ear function the sound is usually perceived as being much louder. Patients with either normal hearing acuity or with a sensorineural hearing loss perceive the tuning fork better by air conduction than by bone conduction. Conversely, patients with conductive losses hear the tuning fork better by bone conduction than air conduction – a negative test result.

**The Weber test**

In this test, the tuning fork is placed in the middle of the forehead or on the vertex, and the patient is asked to signify in which ear they hear the sound louder. In the Weber test, the sound is either heard better by one ear (in other words, it lateralizes), or it is heard equally by both ears. The sound lateralizes to the ear with the better hearing in sensorineural hearing loss, or to the ear with the greater conductive hearing loss in patients with conductive hearing losses. The normal response is to hear the sound equally in both ears, but this result can also be obtained in patients with equal sensorineural or conductive hearing loss.

**The absolute bone-conduction test**

In this test, an assessment is made of the patient’s ability to hear by bone conduction. This is a measure of sensorineural deafness, and the patient’s response is compared with the examiner’s perception of the tone. If the examiner has roughly normal hearing, the patient ought to hear a tuning fork placed on their mastoid as long as the examiner does. If they hear it for less time, it is considered that their bone conduction is diminished and they have worse hearing than the examiner, probably a sensorineural hearing loss.

**AUDIOMETRY**

**Pure-tone audiometry**

A pure-tone audiometer produces tones of varying intensity (0–120 dB) and frequency (125–8000 Hz).
The test is performed with the patient wearing earphones, usually but not always in a sound-proofed environment. Test sounds at different intensities and frequencies are introduced via the earphones, and patients are asked to indicate when they hear the sound. Threshold values at each frequency are determined and plotted on a graph – an audiogram (Fig. D.1). Normal thresholds have been established by the National Physics Laboratory, and young adults should have threshold values across the test range of 0–10 dB, with 0 dB being the sound pressure level considered to be threshold for 18-year-olds.

Bone conduction is considered to be equivalent to cochlear function as it bypasses the middle-ear conduction mechanism. To obtain bone conduction, a vibrating disc is applied to the patient’s mastoid process. The test is then carried out in the same manner as for air conduction, and the auditory thresholds are obtained and plotted on the audiogram. In this way, it can be seen whether bone conduction is better than air conduction (conductive deafness), or if bone conduction and air conduction are roughly equal (normal or sensorineural deafness).

**Speech audiometry**

It is possible for a patient to produce a normal pure-tone audiogram yet be unable to interpret or discriminate speech. A more discriminatory test of hearing is a speech audiogram, where the patient has to respond to a list of test words played at threshold levels through earphones. Either a graph can be made of the patient’s speech responses at each test level, or a simple raw speech discrimination score can be recorded. The threshold level at which approximately 50 per cent of the words are interpreted correctly (the half-peak level) is about equivalent to the pure-tone average. A difference between the half-peak level of a speech audiogram and the pure-tone average audiogram is indicative of malingering. Patients with pure conductive losses achieve 100 per cent discrimination at an elevated sound presentation level, and they sustain this increased sound pressure level. Conversely, patients with sensorineural hearing losses rarely achieve 100 per cent discrimination, and with increased sound pressure levels the scores deteriorate – a phenomenon known as ‘roll-over’.

**Tympometry**

Tympometers measure middle-ear compliance. Sound is injected into the ear canal, and the reflected sound level is measured while the atmospheric pressure of the ear canal is varied. The reciprocal value of the reflected sound – the compliance – is plotted against the atmospheric pressure. The compliance of the middle ear is greatest when the external ear canal pressure is the same as that in the middle ear. This test can diagnose middle-ear effusions, ossicular disruptions or discontinuity, and Eustachian tube dysfunction. Low compliance levels are seen in severe otosclerotic fixation of the stapes footplate. Changes in middle-ear compliance produced by stapedius muscle contraction can also be measured by injecting high-intensity sound (threshold 80 dB) to elicit the stapedial reflex. An absence of the reflex or abnormal decay of reflex activity is of diagnostic importance in otosclerosis, ossicular discontinuity and retrocochlear hearing loss.

**Evoked response audiometry**

Sound transduction into neural activity can be detected throughout the auditory pathway. In the cochlea, contraction of the outer hair cells produces echoes, otoacoustic emissions (OAEs) that can be heard by sensitive microphones; depolarization of the first-order neurones can be recorded by electrocochleography and the passage of neural flux in the auditory pathways documented by brainstem and cortical techniques. All of these techniques have specific indications but find universal use for screening neonates for hearing loss, detecting non-organic hearing loss in medicolegal problems, and investigating patients with endolymphatic hydrops.

**Hearing tests in children**

Deafness should be diagnosed in infants as early as possible. The earlier that deafness is diagnosed, the more chance there is of the child developing normal
language skills and not falling behind their peer group from an educational standpoint. Children at particular risk of hearing loss (e.g. those born to deaf parents), premature births requiring intensive care, and infants with abnormal facies, etc., are tested within days of birth or discharge from hospital. For this subset of children, the presence of transient evoked otoacoustic emissions excludes a significant hearing loss in young children. If OAEs are not detected, it is highly likely that the child has a hearing loss of greater than 25–35 dB, and further investigation and rehabilitation measures should be instigated without delay.

Every child born in the UK has a hearing test by specially trained nurses at 6 weeks and at the end of the first year of life. Where the nurse does not receive an unambiguous response indicating normal hearing from simple clinical tests, the child is referred to a special children’s speech and hearing clinic. There, further clinical tests are undertaken, which may include evoked response audiometry.

If a child is found to be deaf, special education is set in hand at a very early age and amplification devices are fitted to the child, depending on the severity of the deafness. Those found to be profoundly deaf are considered for cochlear implantation.

**CONDUCTIVE DEAFNESS**

The common causes of conductive deafness are listed in Box D.1.

Conductive deafness is often less severe than sensorineural deafness. Complete disruption of the sound-transmitting mechanism imparts a 60 dB hearing loss, while a simple perforation inflicts a hearing loss of 25–30 dB. Many of these conductive hearing losses are amenable to surgical correction (e.g. otosclerosis, middle-ear effusions or perforations), and some can be cured by medication (e.g. acute otitis media). Others can be overcome by hearing aids.

**Congenital syndromes**

There are a number of congenital syndromes associated with deafness, and most are sensorineural in nature. Where there is a conductive element to the hearing loss, it is often associated with a cochlear abnormality as well as resulting in a mixed hearing loss. It is quite impossible to list all the possible combinations of congenital abnormalities, but the golden rule is that, if one abnormality is observed, a careful search for others must be made and that, during this search, deafness must never be forgotten. Some of the more common congenital syndromes associated with hearing loss are listed below.

**Goldenhar’s syndrome**

The anomalies characteristically present in this maldevelopment of the first and second branchial arches are microtia, total atresia of the external auditory canal, ossicular abnormalities, absent middle-ear muscles and anomalous facial nerve pathway. Abnormalities of the inner ear may be present, as well as hemifacial microsomia (Fig. D.2).

**Treacher Collins syndrome**

This comprises micrognathia, depressed malar bones, eyes sloping downwards and outwards with notched lower lids, ptosis of the auricles and middle-ear abnormalities with deformed ossicles (Fig. D.3).

**Crouzon’s deformity**

Crouzon’s syndrome is a craniofacial dysostosis characterized by exophthalmos, a divergent squint, hypoplastic maxillae, a short upper lip, hypertelorism, a beak-shaped nose and deafness caused by atresia and middle-ear abnormalities.

**Marfan’s syndrome**

This comprises an inherited collagen disorder, abnormally long extremities, subluxation of the lens, cardiovascular abnormalities and deafness. The auricles are very large in this condition, and the cartilaginous canals tend to collapse.

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<table>
<thead>
<tr>
<th>Box D.1 The causes of conductive deafness</th>
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<tbody>
<tr>
<td><strong>Congenital lesions</strong></td>
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<tr>
<td>• Atresia of the external meatus and middle ear usually with microtia (see Fig. D.2)</td>
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<tr>
<td>• Atresia associated with other facial defects</td>
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<tr>
<td>• Middle-ear deformities</td>
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<tr>
<td>– Some syndromes (frequently associated with sensorineural loss in addition to the conductive loss)</td>
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<tr>
<td>– Mandibulofacial dysostosis (Treacher Collins syndrome)</td>
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<tr>
<td>– Crouzon’s deformity</td>
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<td>– Marfan’s syndrome</td>
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<tr>
<td>– Klippel–Feil syndrome</td>
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<tr>
<td>– Trisomy D and E</td>
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<tr>
<td>– Cretinism</td>
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<tr>
<td>– Cleft palate</td>
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<tr>
<td>– Submucous cleft palate</td>
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<tr>
<td>– Osteogenesis imperfecta (van der Hoeve–de Kleyn triad)</td>
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<tr>
<td><strong>Middle-ear lesions</strong></td>
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<tr>
<td>• Trauma</td>
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<tr>
<td>• Blood</td>
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<tr>
<td>• Oscillar disruption</td>
</tr>
<tr>
<td>• Perforated tympanic membrane</td>
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<tr>
<td>• Acute otitis media</td>
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<tr>
<td>• Eustachian malfunction</td>
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<tr>
<td>– Atelectasis of middle ear</td>
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<tr>
<td>– Serous otitis (‘glue ear’)</td>
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<tr>
<td>• Otitic barotrauma</td>
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<tr>
<td>• Chronic otitis media</td>
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<tr>
<td>• Haemotympanum</td>
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<tr>
<td>• Malignant disease</td>
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<tr>
<td>• Glomus tumour</td>
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<tr>
<td>• Otosclerosis</td>
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Klippel–Feil syndrome
This is a syndrome in which there are malformed cervical vertebrae and a webbed neck in association with hearing loss caused by ankylosis of the ossicles. Some patients also have rudimentary inner ears.

Trisomy D and E
Patients with this condition have low-set ears, preauricular tags, atresia of the external auditory canals and an absence of the middle-ear cleft.

Osteogenesis imperfecta
Sometimes known as the van der Hoeve’s syndrome, the classical triad is deafness caused by stapedial fixation and incudo-stapedial fragility, blue sclera and fragile bones. It is fortunately an uncommon condition, with a frequency of 2–3 per 100 000 population.

Cleft palate
Cleft palate is one of the most common congenital deformities, with a frequency of about 1 in 1000. It is also a cause of deafness. The palatal muscles (tensor palati and levator palati) play an important part in Eustachian tube opening and closure. Nearly every child with a cleft palate has a middle-ear effusion, and some go on to develop atelectasis of the middle-ear cleft. Closure of the palatal defect does not influence Eustachian tube function. Submucous clefts of the palate in which there is a deficiency of the muscular layer are equally disruptive to Eustachian tube function. Some of these children are helped by reconstructive surgery. This can be a difficult condition to recognize but, in a few, a bifid uvula may be present and draw attention to the abnormality (Fig. D.4).

Figure D.2 Atresia of the right ear, showing an absent external auditory meatus and deformed auricle.

Figure D.3 Treacher–Collins syndrome, showing the typical appearance of the eye, micrognathia, depressed malar bone and ptosis of the ear.

Figure D.4 Submucous cleft palate with the deeply bifid uvula typical of this rare condition.
Other syndromes associated with hearing loss

The hearing losses found in these syndromes are often predominantly sensorineural, with relatively minor conductive elements. Examples are Pendred’s syndrome, in which hearing loss is associated with thyroid deficiency, rubella infection and thalidomide toxicity.

Abnormalities of the external auditory canal

Diseases of the external auditory meatus rarely cause deafness, as hearing is retained while there is still the smallest airway past the obstruction to the tympanic membrane. However, sudden deafness can develop when wax becomes impacted or wet, when it expands and closes the canal. Similarly, the oedema associated with otitis externa can also cause occlusion of the ear canal and a conductive hearing loss.

Middle-ear causes of hearing loss

Most of these are easily identified by a thorough and careful otoscopic examination.

Otitis media with effusion (secretory otitis media, ‘glue ear’)

This is an extremely common condition and is present in 4 per cent of all children aged between 5 and 15 years. This means that almost every classroom in the country will contain one child with a temporary hearing loss caused by ‘glue ear’. The resulting hearing loss is variable but may be sufficient to impair education. Fairly typical changes occur in the eardrum, which becomes retracted and develops a yellowish glaze. Occasionally, fluid levels and bubbles can be seen through the eardrum, but in other instances there may be no observable clinical signs. The diagnosis is made by tympanometry, where a flat compliance curve is recorded. In adults, otitis media with effusion may be the first sign of Eustachian tube obstruction by a nasopharyngeal carcinoma. Although less common among Europeans, this condition is of very definite clinical significance in the Chinese, where nasopharyngeal carcinoma is the most common head and neck tumour. Drainage of the effusions and the insertion of grommets (ventilation tubes) restores hearing.

Acute otitis media

This is a common viral or bacterial infective disorder of childhood caused by Streptococcus pneumoniae, Haemophilus influenzae, Moraxella catarrhalis and, less frequently, Streptococcus pyogenes or Staphylococcus aureus. Acute otitis media often accompanies or complicates upper respiratory tract infections. The child becomes unwell and pyrexic, and complains of a hearing loss in the affected ear and increasing earache. Occasionally, instead of complaining of earache, a child may complain of abdominal pain. Eventually, the eardrum ruptures to release pus that has accumulated in the middle-ear cleft. Pain diminishes rapidly once the pus has drained, as do any toxic symptoms and signs. Over the course of the next few days, the infection resolves and the ruptured eardrum seals spontaneously in most cases. There is a considerable dichotomy of opinion about the value of antibiotics in the management of this condition. One body of opinion suggests that it can be managed just as successfully by symptomatic treatment with analgesics, while others feel that antibiotics should always be prescribed.

Chronic otitis media

Two types of chronic otitis media are recognized, tubo-tympanic and attico-antral disease. Tubo-tympanic disease is characterized by a persistent central perforation, while attico-antral disease is associated with ingrowth of skin from the attic or postero-marginal region of the tympanic membrane into the mastoid antrum, mastoid and middle ear – cholesteatoma. Both groups of patients develop hearing loss in the affected ear and are subject to chronic discharge. In the case of tubo-tympanic disease, this discharge is mucoid and not particularly offensive. By contrast, patients with atticoantral disease have a watery and offensive discharge.

Both conditions are potentially dangerous, as uncontrolled infection may spread to cause meningitis, brain abscess and facial palsy. Persistent perforations can often be repaired surgically, but those which cannot should be kept dry and free from water contamination. Episodes of infection are treated with topical antibiotics. Attico-antral disease almost always requires surgical treatment in the form of a mastoid exploration. Some small cholesteatomas can be managed by suction clearance on an intermittent basis, but these cases are relatively rare.

Otosclerosis

This is a form of deafness caused by fixation of the stapes by the development of new bone around its footplate. It tends to affect young adults and is a progressive form of deafness that increases as the footplate becomes more and more fixed. In females, the deafness is often made worse by pregnancy. Pure-tone audiometry demonstrates a conductive loss often with an increased loss at 2 kHz, the so-called ‘Carhart’s notch’. Treatment of this condition is
DEAFNESS

determined by the degree of hearing loss it inflicts, the likely benefit derivable from surgical intervention and the patient’s wishes. Oral fluoride therapy can retard progression. Those with significant hearing loss will certainly benefit from a hearing aid, and some from removal of the stapes and replacement by a prosthesis (stapedectomy).

SENSORINEURAL DEAFNESS

Although there may be some overlap, it is probably better to separate those conditions which generally arise in childhood (Box D.2) from those which develop in adult life. The importance of early diagnosis of hearing deficit in childhood has already been mentioned. The normal development of an infant is greatly dependent upon hearing, the understanding of speech being the one function of human behaviour that sets man apart from animals. Failure to hear speech not only prevents the development of language, but it also inhibits the formation of personal and social relationships. Much can be achieved nowadays to minimize the handicap of hearing loss by cochlear implantation or the provision of suitable hearing aids and peripatetic care.

Infants who have a family history of deafness, maternal infection during pregnancy or perinatal problems, who are late to talk or who have other congenital defects must be considered as being ‘at risk’ of a significant hearing deficit and should be carefully tested. The frequency of sporadic cases of deafness makes testing of all babies important. In this respect, the mother’s views should never be ignored; if she thinks her child is deaf, the diagnosis should be presumed correct until demonstrated to be otherwise. It must be remembered also that mild or moderate conductive deafness may be an additional handicap in this age group. This added handicap may be the decisive factor that prevents a child from hearing at all without amplification.

Genetically determined syndromes

There are a very large number of genetically determined syndromes that include sensorineural hearing loss. The most common have been listed together here with a brief description of their main characteristics.

Four main types of structural abnormality associated with severe hearing loss have been described:

- **The Scheibe structural abnormality** is characterized by a failure of sacculo-cochlear development.
- **In the Mondini type of dysplasia**, both vestibular and cochlear structures are deformed.
- **In the Bing abnormality**, there is a normal bony labyrinth but the membranous labyrinth is malformed or degenerate in both the cochlea and vestibule. In children with this malformation, other central nervous system abnormalities may be present.
- **The Michel group of abnormalities mainly affects the otic capsule with almost complete lack of development of the inner ear**. This particular type of abnormality is often associated with mental deficiency.

Non-genetically determined syndromes

These include the following:

- **Waardenburg’s syndrome**: this is an example of the group of integumentary system disease and deafness. A white forelock and heterochromia of the iris are combined with familial genetic deafness.
- **Pendred’s syndrome**: this is a syndrome that comprises congenital goitre and hypothyroidism with severe abnormalities of the labyrinth in both the vestibular and cochlear parts. A large group of abnormalities is described in which hearing loss is associated with eye disease, retinal abnormalities, myopia, optic atrophy and corneal degeneration. For example, **Usher’s syndrome** is deafness combined with retinitis pigmentosa.

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**Box D.2 Causes of deafness in children**

<table>
<thead>
<tr>
<th>Prenatal</th>
<th>Perinatal</th>
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<tbody>
<tr>
<td>Genetic</td>
<td>Genetic</td>
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<tr>
<td>• Scheibe type</td>
<td>• Prematurity</td>
</tr>
<tr>
<td>• Bing–Siebenmann type</td>
<td>• Jaundice – haemolytic disease and kernicterus</td>
</tr>
<tr>
<td>• Waardenburg’s syndrome</td>
<td>• Anoxia due to birth trauma</td>
</tr>
<tr>
<td>• Pendred’s syndrome</td>
<td>Postnatal</td>
</tr>
<tr>
<td>• Mondini–Alexander type</td>
<td>Genetic</td>
</tr>
<tr>
<td>• Michel type</td>
<td>• Familial degenerative deafness</td>
</tr>
<tr>
<td>• Usher’s syndrome</td>
<td>• Otosclerosis</td>
</tr>
<tr>
<td>• Endemic cretinism</td>
<td>• Alport’s syndrome</td>
</tr>
<tr>
<td>• Klippel–Feil syndrome</td>
<td>Non-genetic infectious diseases</td>
</tr>
<tr>
<td><strong>Non-genetic</strong></td>
<td>• Measles</td>
</tr>
<tr>
<td>• Diseases occurring in pregnancy</td>
<td>• Mumps</td>
</tr>
<tr>
<td>– Rubella and other viral illnesses</td>
<td>• Meningitis</td>
</tr>
<tr>
<td>– Toxaemia</td>
<td>– Meningococcal</td>
</tr>
<tr>
<td>– Diabetes</td>
<td>– Pneumococcal</td>
</tr>
<tr>
<td>– Syphilis</td>
<td>– Viral</td>
</tr>
<tr>
<td>– Nephritis</td>
<td>– Tuberculous</td>
</tr>
<tr>
<td>• Drugs taken in pregnancy</td>
<td>– Trauma</td>
</tr>
<tr>
<td>– Streptomycin</td>
<td>– Otitis media</td>
</tr>
<tr>
<td>– Quinine</td>
<td>• Otoxic antibiotics</td>
</tr>
<tr>
<td>– Salicylates</td>
<td>– Streptomycin</td>
</tr>
<tr>
<td>– Thalidomide</td>
<td>– Neomycin</td>
</tr>
<tr>
<td></td>
<td>– Gantamicin</td>
</tr>
</tbody>
</table>
Late-onset genetic deafness is now considered to be an important cause of deafness in childhood and later life than had been previously thought. Alport’s syndrome is an example of this and is also representative of a group of syndromes where genetic hearing loss is associated with nephritis and lenticonus. Non-genetic prenatal influences are also well recognized as causes of significant hearing loss. Rubella is the most widely known example and one that is potentially preventable by the use of vaccination programmes. Other perinatal causes of hearing loss, for example perinatal anoxia or jaundice, are also potentially preventable.

Postnatal causes of profound sensorineural hearing loss include mumps and meningitis. Curiously, the hearing loss acquired with mumps is nearly always unilateral, and this serves to identify it on occasion.

The proportion of the various groups of conditions causing congenital deafness has been estimated as one-quarter each of genetic, maternal rubella, perinatal causes and unknown. Genetic causes — either alone or as a contributory factor by increasing the liability to other influences — are being increasingly implicated. In areas of the world where consanguineous marriages are common, genetic sensorineural deafness is reported to comprise 70–80 per cent of cases. A careful family history is most important in making a probable diagnosis.

A different scheme of classification is appropriate for adults (Box D.3).

More common causes of sensorineural deafness

Many of these conditions in adults have been considered above. Some, however, require further attention.

Refsum’s disease is yet another syndrome combining eye disease (retinitis pigmentosa), polyneuritis, cerebellar ataxia and a genetic late-onset deafness. The inheritance is autosomal recessive.

Acoustic trauma from noise is wholly preventable and entirely untreatable. It may be diagnosed from the history and an audiogram that shows a typical curve, sharply falling in the higher frequencies with a characteristic dip at 4 kHz (Fig. D.5). Noise-induced hearing loss may be demonstrated audiometrically as either temporary (noise-induced temporary threshold shift) or permanent (noise-induced permanent threshold shift), and it may be caused by sudden loud sounds such as gunfire or by continuous trauma such as traffic noise, industrial noise, agricultural noise or even pop music. Compensation is awarded on a very large scale to workers exposed to noise in industry who have not been provided with and made to use noise protection. This affects mainly shipyard and railway workers, motor car industry workers and miners. Unfortunately, no amount of compensation can make up for the hearing loss or impaired quality of life. Tinnitus is very frequently present in noise-induced hearing loss and contributes to the misery experienced by those deafened by this means.

Vascular lesions of the inner ear are the cause (or part cause) of many cases of deafness. Sudden, small vascular accidents in the end-arterioles may cause deafness by damaging part or the whole of the organ of Corti.

---

**Box D.3 Causes of deafness in older children and in adults**

<table>
<thead>
<tr>
<th>Cochlear lesions</th>
<th>Retrocochlear lesions</th>
</tr>
</thead>
<tbody>
<tr>
<td>● Late-onset genetic deafness</td>
<td>● Drug-induced deafness</td>
</tr>
<tr>
<td>○ Familial degenerative</td>
<td>○ Antibiotics</td>
</tr>
<tr>
<td>○ Alport’s syndrome</td>
<td>○ Aminoglycosides: streptomycin, neomycin, gentamicin</td>
</tr>
<tr>
<td>○ Refsum’s syndrome</td>
<td>○ Others in large doses</td>
</tr>
<tr>
<td>○ Otosclerosis (later cochlear effects)</td>
<td>○ Aspirin (reversible deafness)</td>
</tr>
<tr>
<td>○ Inflammatory (labyrinthitis)</td>
<td>○ Quinine</td>
</tr>
<tr>
<td>○ Bacterial</td>
<td>○ Chloroquine</td>
</tr>
<tr>
<td>○ Late-onset rubella</td>
<td>○ Chemotherapeutic agents for malignant disease</td>
</tr>
<tr>
<td>○ Syphilis</td>
<td>○ Unknown</td>
</tr>
<tr>
<td>○ Mumps</td>
<td>● Meningitis</td>
</tr>
<tr>
<td>○ Herpes</td>
<td>● Leptomenigitis</td>
</tr>
<tr>
<td>○ Measles</td>
<td>● Syphilitic</td>
</tr>
<tr>
<td>○ Trauma</td>
<td>● Tuberculous</td>
</tr>
<tr>
<td>− Fracture of temporal bone</td>
<td>● Cerebellar ataxic neuropathy</td>
</tr>
<tr>
<td>− Acoustic trauma</td>
<td>● Vogt-Koyanagi syndrome</td>
</tr>
<tr>
<td>○ Temporary</td>
<td>● Harada’s disease</td>
</tr>
<tr>
<td>○ Permanent</td>
<td>● Unknown</td>
</tr>
<tr>
<td>● Vascular lesions</td>
<td>● Central</td>
</tr>
<tr>
<td>○ Atherosclerosis</td>
<td>● Multiple sclerosis</td>
</tr>
<tr>
<td>○ Hypertension</td>
<td>● Encephalitis</td>
</tr>
<tr>
<td>○ Vascular accident of end-artery</td>
<td>● Meningomyelitis</td>
</tr>
<tr>
<td>● Ménière’s disease</td>
<td>● Pontine glioma</td>
</tr>
<tr>
<td>(labyrinthine hydrops)</td>
<td>● Concussion</td>
</tr>
<tr>
<td>− Lermoyez’s syndrome</td>
<td>● Vascular accidents</td>
</tr>
<tr>
<td>− Leukaemia</td>
<td>● Brainstem damage from head injury</td>
</tr>
<tr>
<td>− Malaria</td>
<td>● Psychogenic – hysterical</td>
</tr>
<tr>
<td>● Degenerative (partly vascular)</td>
<td>● Unknown</td>
</tr>
<tr>
<td>○ Presbyacusis</td>
<td>● Tinnitus</td>
</tr>
<tr>
<td>● Vitamin deficiency</td>
<td>● Drug-induced deafness</td>
</tr>
<tr>
<td>● Vitamin B deficiency</td>
<td>● Acoustic neuroma (VIIIth nerve neurilemmoma)</td>
</tr>
<tr>
<td>● Dietary</td>
<td>● Meningitis</td>
</tr>
<tr>
<td>○ Tropical ataxic neuropathy</td>
<td>● Leptomenigitis</td>
</tr>
<tr>
<td>● Hormonal</td>
<td>● Syphilitic</td>
</tr>
<tr>
<td>− Myxoedema</td>
<td>● Tuberculous</td>
</tr>
<tr>
<td>− Pregnancy</td>
<td>● Cerebellopontine angle tumour</td>
</tr>
<tr>
<td>● Drug-induced deafness</td>
<td>● Trauma</td>
</tr>
<tr>
<td>○ Antibiotics</td>
<td>● Carcinomatous neuropathy</td>
</tr>
<tr>
<td>○ Aminoglycosides: streptomycin, neomycin, gentamicin</td>
<td>● Vogt-Koyanagi syndrome</td>
</tr>
<tr>
<td>○ Others in large doses</td>
<td>● Harada’s disease</td>
</tr>
<tr>
<td>○ Aspirin (reversible deafness)</td>
<td>● Unknown</td>
</tr>
<tr>
<td>○ Quinine</td>
<td>● Central</td>
</tr>
<tr>
<td>○ Chloroquine</td>
<td>● Multiple sclerosis</td>
</tr>
<tr>
<td>○ Chemotherapeutic agents for malignant disease</td>
<td>● Encephalitis</td>
</tr>
<tr>
<td>○ Unknown</td>
<td>● Meningomyelitis</td>
</tr>
<tr>
<td>● Malaria</td>
<td>● Pontine glioma</td>
</tr>
<tr>
<td>● Tinnitus</td>
<td>● Concussion</td>
</tr>
<tr>
<td>● Vascular accidents</td>
<td>● Brainstem damage from head injury</td>
</tr>
<tr>
<td>● Psychogenic – hysterical</td>
<td>● Tinnitus</td>
</tr>
<tr>
<td>● Unknown</td>
<td>● Drug-induced deafness</td>
</tr>
</tbody>
</table>
The cause of Ménière's disease (Fig. D.6) is not fully known and is probably multifactorial, but the end result is an increase in endolymphatic pressure. Episodes of increased pressure give rise to a sensation of fullness in the affected ear that is followed by intense rotatory vertigo and impaired hearing. After a period of hours, the vertigo subsides and hearing improves. With repeated attacks, permanent damage is sustained by both the organ of Corti and the vestibular sensory epithelium. The effect of repeated attacks is cumulative. The diagnosis of Ménière's disease is made from the typical history. The disease shows periods of remission between paroxysms of attacks. Lermoyez's syndrome is a variant of Ménière's disease in which the hearing improves very suddenly after an attack of vertigo and tinnitus. It is thought that the membranous labyrinth ruptures, releasing the endolymphatic pressure and restoring cochlear function. Other conditions can also cause endolymphatic hydrops, for example myxoedema and post-meningitic and head injury syndromes.

Leukaemia causes haemorrhage in the inner ear, whereas in malaria destruction of the blood cells results in pigment being left in the cells. Deafness in this disease may also be caused by antimarial drugs. Presbyacusis (senile deafness) is common to mankind, and loss of hearing in the higher frequencies is almost invariable with age, although the rate is dependent upon genetic background, exposure to noise (city dwellers losing their hearing more rapidly than country dwellers) and vascular changes associated with atherosclerosis. The audiogram in Figure D.7 shows an increasing depression in the high frequencies. Failure to discriminate speech ('I can hear you talking, but I cannot hear what you say'), particularly in background noise, is characteristic of this high-tone hearing loss as the consonant sounds that give speech its intelligibility are carried in this part of the frequency spectrum.

Drug-induced deafness, ototoxicity, not only results from systemic treatment but can also develop following the excessive use of topical eardrops. Aminoglycoside antibiotics, loop diuretics, cytotoxic agents, quinine and aspirin are the most commonly implicated substances. Among the neural and central lesions, vestibular schwannomas (acoustic neuromas) are important because they are silent, slow-growing, difficult to diagnose and potentially lethal if not
found when reasonably small. They are usually solitary but in patients with neurofibromatosis type 2 are characteristically bilateral and associated with other intracranial tumours. The presentation of these tumours is usually with a progressive unilateral sensorineural deafness often accompanied by tinnitus. Vertigo or unsteadiness is also not uncommon and is frequently misdiagnosed as Ménière’s disease. As the tumour becomes larger, trigeminal symptoms arise that include progressive sensory deficits or even trigeminal neuralgia. Diagnosis of acoustic neuroma after clinical, audiometric and vestibular tests is made by magnetic resonance imaging (Fig. D.8).

The Vogt–Koyanagi syndrome, from which it is thought that the artist Goya suffered, is a sudden and rare illness with severe headache and malaise that goes on to uveitis, alopecia, vitiligo and deafness. Harada’s disease is very similar, but with retinal detachment instead of uveitis. The deafness is usually permanent, but the uveitis recovers. The depressing effects of sudden complete deafness on a sensitive artist such as Goya explain his change of style from brightly coloured, happy pictures of handsome men and pretty girls to those of his later ‘Black Period’ and the ‘Disasters of War’. This is an indication of the severe psychological effects that deafness may bring.

A delusion may be defined as a false unshakeable belief that is out of keeping with the patient’s educational, cultural and social background. It is a belief that is held with pathological certainty. Patients themselves may not complain of this symptom but will come to medical attention because they have acted on a delusional belief, or relatives have become aware of the unusual or bizarre content of their ideas. A delusion should be distinguished from an overvalued idea, which is a preoccupying, comprehensible conviction held beyond the bounds of reason. For example, morbid jealousy is an overvalued idea when a spouse who is unduly preoccupied by thoughts of their partner’s suspected infidelity can be reassured after lengthy persuasion that their belief is irrational. In delusional jealousy, on the other hand, such reassurance would not be possible because the spouse would be convinced of infidelity in the face of all evidence to the contrary. Delusions are also to be distinguished from obsessional ideas, which may be bizarre but trouble the patient and are seen by them as intrusive, unwanted and requiring some response (see OBSESSIONS, p. 470). Similarly, religious and political non-conformity, however extreme, does not represent delusional belief when it is in keeping with the culture to which the patient belongs.

Delusions may be caused by organic disease, by drug or substance abuse or may be signs of a psychotic illness (Box D.4).

### ORGANIC DELUSIONAL STATE

Delusions may be a prominent feature of a number of quite diverse conditions, and organic causes should be considered in any patient, particularly if delusions

#### Box D.4 Causes of delusions

<table>
<thead>
<tr>
<th>Common</th>
<th>Prescribed drugs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Schizophrenia</td>
<td>– Bromocriptine</td>
</tr>
<tr>
<td>Psychotic depression</td>
<td>– L-Dopa</td>
</tr>
<tr>
<td>Mania</td>
<td>– Corticosteroids</td>
</tr>
<tr>
<td>Delusional disorders</td>
<td>Alzheimer’s-type dementia</td>
</tr>
<tr>
<td>Substance abuse</td>
<td>Multi-infarct dementia</td>
</tr>
<tr>
<td>– Alcohol</td>
<td>Temporal lobe epilepsy</td>
</tr>
<tr>
<td>– Amphetamines</td>
<td>Hyperthyroidism</td>
</tr>
<tr>
<td>– Methylphenidate</td>
<td>Encephalopathies including HIV</td>
</tr>
<tr>
<td>– Cocaine</td>
<td>Head injury</td>
</tr>
<tr>
<td>– Cannabis</td>
<td>Rarer</td>
</tr>
<tr>
<td>– Hallucinogens (e.g. LSD)</td>
<td>– Neurosyphilis</td>
</tr>
<tr>
<td>– Phencyclidine</td>
<td>– Cerebral abscess</td>
</tr>
<tr>
<td>– Delirium</td>
<td></td>
</tr>
</tbody>
</table>
appear for the first time in someone over the age of 35. Substances such as amphetamines, cocaine and phencyclidine cause an initial feeling of well-being and confidence. Intoxication with high doses may, however, lead to an episode of paranoid delusions, with visual, auditory and tactile hallucinations, incoherent speech and anxious mood. Associated with these mental phenomena are tachycardia, pupillary dilatation, elevated blood pressure, sweating and sometimes nausea. Full recovery is usual within 48 hours, but cessation of regular heavy use may lead to a withdrawal state that again may be associated with paranoid delusions and suicidal ideation, fatigue, depression and agitation persisting for several days. Cocaine, shortly after intake, leads in some users to the rapid onset of a delusional disorder, which can persist for over a week and occasionally for several months. Delusions are usually persecutory, and other features are body image distortion, the feeling of insects on the skin (formication, or the ‘cocaine bug’) and sometimes aggression directed against imagined persecutors. Cannabis and hallucinogens such as LSD may also cause paranoid delusions in some users.

An organic delusional state may develop in some subjects with temporal lobe epilepsy who show interictal features similar to schizophrenia.

DELUSIONS IN THE PSYCHOSES

In the absence of organic causes, delusional thinking is commonly a symptom of schizophrenia, the delusional disorders, mania or psychotic depression. To distinguish these possibilities, considerable stress is placed on whether or not the content of the delusions is congruent or incongruent with the prevailing mood of the patient. For example, when a depressed patient expresses the delusion that there is a plot to kill him, this is in keeping with the depressed mood, as is the case with a manic patient who believes he has been invested with special powers and has a mission to save the world. These delusions are said to be secondary because an understandable connection can be made between the patient’s mood and the content of the delusion. Moreover, these beliefs will disappear when the mania and depression have responded to treatment. In schizophrenia, by contrast, the content of delusional thinking is often bizarre and out of keeping with the patient’s mood state, and the beliefs often persist. Primary delusions describe those beliefs which take the form of sudden convictions that come into the patient’s mind in response to what the ordinary observer would consider a totally unrelated experience, for example if a patient when reading the number plate of a passing car suddenly becomes convinced that the driver was accusing her of being a homosexual. Primary delusions of this sort are observed more often in schizophrenia than in affective illness. Schizophrenic patients may also experience delusional mood, when they are perplexed, unsettled and convinced that something self-referential is occurring but are unable to understand what it is. There is no hard and fast distinction between the types of delusion found in schizophrenia, mania and depression, but clinical observation suggests that certain types of delusion are more commonly associated with schizophrenia. These include the delusion that the body is being influenced by outside forces and the subject is being ‘made’ to perform certain acts, think certain thoughts or, rarely, feel certain emotions. The patient may complain that alien thoughts have been inserted into his head or that his own thoughts have been removed. Some commonly encountered delusions that may occur just as frequently in schizophrenia as affective illness include persecutory beliefs, grandiose delusions, delusions of guilt, poverty and worthlessness, and hypochondriacal beliefs. Both grandiose and depressive delusions are often set in a religious context, although beliefs in influence by radio, radar and other real or imagined physical forces are also common.

Monosymptomatic delusions sometimes occur in a patient whose thinking, mood and behaviour show none of the disturbances normally associated with schizophrenia or affective psychosis. Morbid jealousy, the total conviction of the spouse’s infidelity, although sometimes associated with the psychoses, alcohol abuse or psychopathic personality, may also occur as the only symptom, when it must nevertheless be taken seriously by the clinician as a significant cause of domestic violence and even homicide. Other delusions that occur in isolation as well as in the context of major psychotic illness include delusions of love (de Clerambault’s syndrome) in which the patient believes that some person, usually an authority figure, is in love with them, and delusions of misidentification (Capgras syndrome), in which the patient believes that someone close to them, for example a wife, has been replaced by an imposter pretending to be that person. Somatic (hypochondriacal) delusions take several forms. The person may be convinced that certain parts of the body (brain, intestine or stomach) are not functioning or, in extreme cases, are not there at all. There may be conviction of an internal parasite or infestation on or in the skin. The body may be held to be misshapen or ugly, or there may be a belief that a foul smell is emanating from it. Hypochondriacal delusions are frequently found in psychotic depression,
and they may include the belief that the patient is dying from an incurable disease such as cancer. However, such beliefs may also occur in the absence of other overt psychiatric abnormality. Delusions concerning the face, mouth, teeth and gums may be accompanied by frequent and persistent demands for medical and dental investigations and be the basis for litigation. Some patients with monosymptomatic delusions are considered depressed or schizophrenic; in others, the condition is ill understood and difficult to treat.

Delusions on their own have little diagnostic specificity and their assessment must take account of all aspects of the patient’s physical and mental state as well as their cultural background. Brief delusional ideas may develop in normal people following severe sleep deprivation, and delusions, often of a persecutory nature, will develop in a person in strange or unusual surroundings especially when under stress. In delirium, delusions develop when consciousness is clouded and the cause of the condition may be apparent. Delusions in the delirious (acutely confused) patient tend to be fleeting, poorly formed and drawn from the immediate environment (e.g. ‘the nurses are trying to poison me’). Similarly, in chronic organic states, delusions will rarely be the only presenting feature, and the diagnosis will be suggested by the memory disturbance and other cognitive deficits. The differential diagnosis of substance abuse, schizophrenia and mania gives most difficulty particularly because drug abuse is common in patients with functional psychosis and may be the precipitant as much as the cause of the delusional state (see also HALLUCINATIONS, p. 253). A careful history and urine drug screen should elucidate the role of stimulants and hallucinogens in a psychotic episode.

DEPERSONALIZATION

Andrew Hodgkiss

Depersonalization is the experience of losing the accustomed sense of one’s own reality and should be distinguished from the very much rarer delusion of not being real that is occasionally elicited in schizophrenia. There are a variety of unusual phenomena that can accompany depersonalization:

- **Derealization** (a sensation that the outer world is not real)
- **Emotional numbing**
- **Bodily change** (particularly enlargement of the head or limbs, lifelessness and unfamiliarity).
- **Autoscopy** (seeing yourself from a distance)
- **Doubling** (perceptions linked to an independently existing double)

- **Automaton experience**, with the self or others appearing to act or think in a contrived, forced fashion
- **Disturbance of time sense**
- **Dizziness**, which is a particularly common associated symptom

Typically, depersonalization is alarming and distressing, which contrasts sharply with the subjective inability to feel emotions: insight is invariably retained (Box D.5). Depersonalization can be a normal experience, occurring in 30–70 per cent of the population. It is most common in late adolescence and early adulthood, when it is usually mild, transient, associated with fatigue and of no clinical significance. Depersonalization also occurs physiologically in the rare states of sleep deprivation, sensory deprivation and the near-death experience.

Depersonalization can be generated by the mental mechanism of dissociation as an anxiety-reducing response in stress; hence, **dissociative depersonalization** tends to be less unpleasant or alarming and not a focus for complaint. This process permits a temporary emotional respite and better coping in circumstances like battle, accident, admission to hospital or appearance in court, and during the early stages of grief. It can also precede anticipated trauma, and indeed in deliberate self-injury (especially multiple cutting) may even facilitate the act through a diminution of pain. If the stress is prolonged,

<table>
<thead>
<tr>
<th>Box D.5 Causes of depersonalization</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Most common</strong></td>
</tr>
<tr>
<td>1. Normal</td>
</tr>
<tr>
<td>2. Stress reaction (dissociative depersonalization)</td>
</tr>
<tr>
<td>3. Anxiety</td>
</tr>
<tr>
<td><strong>Less common</strong></td>
</tr>
<tr>
<td>4. Drugs</td>
</tr>
<tr>
<td>4.1 Alcohol</td>
</tr>
<tr>
<td>4.2 Hallucinogens</td>
</tr>
<tr>
<td>4.3 Stimulants</td>
</tr>
<tr>
<td>4.4 Cannabis</td>
</tr>
<tr>
<td>4.5 Anticholinergics</td>
</tr>
<tr>
<td>4.6 Benzodiazepines</td>
</tr>
<tr>
<td>5. Drug withdrawal</td>
</tr>
<tr>
<td>6. Fever</td>
</tr>
<tr>
<td>7. Neurological disorders</td>
</tr>
<tr>
<td>7.1 Head injury</td>
</tr>
<tr>
<td>7.2 Brain tumour</td>
</tr>
<tr>
<td>7.3 Encephalitis</td>
</tr>
<tr>
<td>7.4 Multiple sclerosis</td>
</tr>
<tr>
<td>7.5 Epilepsy</td>
</tr>
<tr>
<td>8. Psychiatric disorders</td>
</tr>
<tr>
<td>8.1 Phobia</td>
</tr>
<tr>
<td>8.2 Panic</td>
</tr>
<tr>
<td>8.3 Depression</td>
</tr>
<tr>
<td>8.4 Obsessive–compulsive disorder</td>
</tr>
<tr>
<td>8.5 Schizophrenia</td>
</tr>
<tr>
<td>8.6 Post-traumatic stress disorder</td>
</tr>
<tr>
<td>9. Insomnia</td>
</tr>
<tr>
<td>10. Primary depersonalization</td>
</tr>
<tr>
<td>11. Syndrome</td>
</tr>
<tr>
<td><strong>Rare</strong></td>
</tr>
<tr>
<td>12. Psychiatric disorders</td>
</tr>
<tr>
<td>12.1 Hysteria</td>
</tr>
<tr>
<td>12.2 Malingering</td>
</tr>
<tr>
<td>12.3 Sleep deprivation</td>
</tr>
<tr>
<td>12.4 Sensory deprivation</td>
</tr>
<tr>
<td>12.5 Near-death experience</td>
</tr>
<tr>
<td>12.6 Brainwashing</td>
</tr>
</tbody>
</table>
depressive depersonalization can persist equally long, for months or even years, as recounted in the experiences of concentration camp survivors.

Depersonalization in the setting of psychological disorders is also common, but it is unusual for this feature to persist or to be the main complaint. The most common primary conditions are anxiety, agoraphobia and depression. In anxiety, agoraphobia, panic disorder and chronic hyperventilation syndrome, depersonalization is rarely intense or accompanied by associated experiences apart from derealization, while the amount of anguish and self-concern is substantial, with the fear of going mad frequently expressed. In depression and post-traumatic stress disorder, emotional numbing characteristically accompanies depersonalization, and it can even be the predominant feature – bodily change, especially physical lifelessness, may be reported in severe depression. In schizophrenia, depersonalization is usually an early, transient phenomenon, with more typical disturbances of identity either present or emergent soon afterwards – autoscopy, often with emotional detachment, tends to occur when the illness is acute and established.

Drug intoxication or withdrawal can precipitate depersonalization, the most common associations being reported with LSD, mescaline, cannabis and anticholinergic agents. Brain disease or damage may precipitate or even present with depersonalization. Post-concussion syndrome is usually apparent from the history and accompanying features. However, in temporal lobe disease and/or epilepsy, depersonalization may not only be the prime symptom but can also precede positive findings upon neurological examination and investigation. This possibility ought to remain under consideration when the state of depersonalization is severe and persistent, autoscopy or perceptual distortions are present, and there are no apparent stressors or evidence of a primary emotional illness.

Primary depersonalization syndrome is an uncommon condition that presents abruptly in early adulthood with recurrent bouts of severe depersonalization, usually with many of the concomitant features. There are characteristic personality traits associated with this condition, the typical sufferer being described as intelligent, obsessive, introspective, sensitive and hypochondriacal. These patients often have particular difficulty in establishing personal relationships and coping with the general upheaval of adolescence. Their disorder tends to be chronic, readily precipitated by stress and anxiety, but with minimal impairment in social functioning; they are prone to hypochondriasis in later life.

DEPRESSION

Andrew Hodgkiss

Depressed mood is one of the most common conditions for which patients seek help. Unhappiness is an understandable response to loss of any sort. Depression refers to a more severe, and qualitatively different, mood change, when the patient may describe feelings of hopelessness, helplessness, worthlessness, guilt, suicidality, tiredness and fatigue. However, not infrequently, one has to rely on a history from the patient, their friends and employers to recognize severe depressive illness in some people who do not themselves acknowledge mood change (smiling or masked depression). For cultural and other reasons, some patients have no words for moods (alexithymia), and may communicate their distress by hypochondriacal complaints or by seeking advice for marital, employment or a variety of other difficulties.

In uncomplicated bereavement, a patient responds to loss by an immediate experience of shock, numbness and feelings of emptiness. There is an initial tendency to deny that the loss has occurred, but this gives way to a depressed mood, often with symptoms of anxiety and panic. There may be poor appetite, with weight loss, insomnia, impaired concentration and feelings of guilt surrounding the death. Auditory and visual hallucinations of the deceased are reported in 10 per cent of instances. The state of grief is resolved as the person accepts the loss and begins to redirect their life and activities. Many cases of depression follow a real or intrapsychic loss event (such as the fall of a cherished ideal or the abandonment of a long-held ambition), and symptoms are not dissimilar to those of bereavement. However, major depressive illness may develop without any clear-cut relation to loss, and classification is based on symptoms, severity and outcome without making any aetiological assumptions. In major depression, there is a distinct quality of depressed mood that is perceived by the patient as being distinctly different from the feelings of loss experienced in bereavement. These patients describe a loss of pleasure in all their activities (anhedonia), and they cannot be even temporarily cheered out of their depression by something good happening in their life. This type of depressive illness is often associated with diurnal variation of mood (the depression being worse in the morning), early-morning waking, psychomotor retardation or agitation, significant appetite impairment and
excessive guilt. In the most severe forms, the patient may express delusions of poverty, ill-health and persecution congruent with depressed mood, and hallucinations, which are often voices critical of the patient. When delusions or hallucinations figure, we use the term psychotic depression. Psychotic depression is more common in older adults than younger. There is a high risk of suicide, and suicidal ideation must always be specifically asked about. There is frequently a family history of depression, and although the illness may develop suddenly, without a recognizable precipitant, adverse stressful life events will often have occurred during the weeks preceding the onset of the illness.

Major depressive illness can be usefully subdivided into bipolar depression (patients have at some stage in their lives experienced an episode of mania) and unipolar depression (patients have repeated bouts of depression without mania). Dysthymia is a chronic depressed mood persisting for most of the time for 2 years or more. Such patients may present with an inability to cope with everyday life, low self-esteem, tiredness, sleep disturbance, somatic complaints and frequently symptoms of anxiety.

Difficulties may arise in the diagnosis of depression, especially when a patient presents with physical symptoms such as headache, low back pain, weight loss, constipation, loss of libido, loss of energy or anorexia. Some patients are worried that they may have cancer, a sexually transmitted disease, angina or memory loss, and the physician’s attention is directed towards an investigation of these physical conditions. In the elderly, depression may present as a pseudodementia that may be initially hard to distinguish from dementia, but which should respond well to standard antidepressant measures.

Anxiety may be a principal feature of depressive illness, and such cases may be termed agitated depression. The patient may be restless and unable to relax during the interview: wringing hands, pulling hair, shifting legs or pacing around the room. Obsessional symptoms can develop during a depressive illness, and these symptoms will generally improve when the depression is treated. However, patients with obsessional personality traits or obsessive–compulsive disorder are also very prone to develop depressive episodes.

DEPRESSION IN OTHER PSYCHIATRIC ILLNESSES
Depression is a common accompaniment of all the other psychoses. An episode of depression may herald the onset of a schizophrenic illness, and patients with schizophrenia are more prone to suffer from depression, perhaps because of the nature of their symptoms, the social and other handicaps of the illness, or perhaps because mood change is an integral part of the disease itself. Depression is particularly common and difficult to treat immediately following the resolution of an acute exacerbation of schizophrenia – post-psychotic depression.

PHYSICAL DISEASE AND DEPRESSION
A number of possible mechanisms account for the very high instance of depression in almost all types of physical disease (Box D.6). Pain, incapacity and loss of health, independence and social status are entirely understandable causes of depression during and following illness. Some illnesses, however, are more specifically linked with depression. Several endocrine disorders may present as depression, including hypothyroidism, hypo- and hyperadrenocorticalism, hypoparathyroidism and hypopituitarism. Certain forms of carcinoma, particularly those of the pancreas, the thyroid and the lung, may cause depression months before other symptoms of the tumour are manifest, possibly due to brain

<table>
<thead>
<tr>
<th>Box D.6 Physical causes of depression</th>
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<tbody>
<tr>
<td><strong>Physical diseases that may cause depression</strong></td>
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<tr>
<td>Infections</td>
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<tr>
<td>• Influenza</td>
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<td>• Hepatitis</td>
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<td>• Infectious mononucleosis</td>
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<td>• Brucellosis</td>
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<td>• Toxoplasmosis</td>
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<tr>
<td>Endocrine disorders</td>
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<td>• Hypothyroidism</td>
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<tr>
<td>• Hyperadrenocorticalism</td>
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<tr>
<td>• Hypopituitarism</td>
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<tr>
<td>• Pancreatic, lung, thyroid carcinomas</td>
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<tr>
<td>• Acute intermittent porphyria</td>
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<tr>
<td>• Systemic lupus erythematosus</td>
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<tr>
<td>• Folate or B₁₂ deficiency</td>
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<tr>
<td>Neurological diseases</td>
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<tr>
<td>• Multi-infarct dementia</td>
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<td>• Alzheimer’s dementia</td>
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<tr>
<td>• Frontotemporal dementias, e.g. Pick’s disease</td>
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<td>• Huntington’s disease</td>
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<tr>
<td>• Multiple sclerosis</td>
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<td>• Parkinson’s disease</td>
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<tr>
<td>• Temporal lobe epilepsy</td>
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<tr>
<td>• Head injury</td>
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<tr>
<td>• Cerebrovascular disease and stroke</td>
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<tr>
<td>• Subdural haematoma</td>
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<td>• Cerebral tumour</td>
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<td>• Encephalitis</td>
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<tr>
<td>• Neurosis</td>
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<tr>
<td>• AIDS</td>
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<tr>
<td><strong>Drugs that can cause depression</strong></td>
</tr>
<tr>
<td>• Reserpine</td>
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<tr>
<td>• Alpha-methyl-dopa</td>
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<td>• Beta-blockers</td>
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<td>• Corticosteroids</td>
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<td>• Oral contraceptives</td>
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<td>• L-Dopa</td>
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<td>• Indometacin</td>
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<td>• Isoniazid</td>
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<tr>
<td>• Cycloserine</td>
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<tr>
<td>• Withdrawal from alcohol</td>
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<tr>
<td>• Withdrawal from amphetamine and related drugs (fenfluramine, diethylpropion)</td>
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<tr>
<td>• Withdrawal from cocaine and phencyclidine</td>
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<tr>
<td>• Use of hallucinogens</td>
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<tr>
<td>• Opioid intoxication</td>
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</table>
active peptides synthesized and released from the tumour. With most central nervous system diseases, depression is a common accompaniment. In multi-infarct and Alzheimer’s-type dementia, depressive symptoms may arise because the patient is aware of their failing powers, but direct cerebral damage may also be implicated. An interaction between a normal response to incapacitating illness and the direct effects of brain damage probably account for the mood disturbances found in Huntington’s disease, multiple sclerosis, Parkinson’s disease, temporal lobe epilepsy, certain types of head injury and cerebrovascular disease.

Drugs that can cause depression
These include both therapeutic agents and drugs of abuse (Box D.6). The former include antihypertensive agents, steroids and antibiotics. Withdrawal from alcohol is a very common cause of depression; withdrawal from amphetamines, the appetite suppressants fenfluramine and diethylpropion, cocaine and phencyclidine may precipitate mood change, particularly in chronic users. Intoxication with opioids and the use of hallucinogens may trigger severe mood disturbance.

Depression may present at any age, and the diagnosis in children is more difficult. In prepubertal children, there may be somatic complaints, agitation, anxiety disorders, avoidance behaviour and phobias. In adolescents, a mood disturbance may be accompanied by negativistic or antisocial behaviour, abuse of alcohol and illicit drugs, feelings of restlessness, aggression, withdrawal from social activities, poor school performance and complaints of not being understood.

DIARRHOEA
Simon Anderson
Diarrhoea is defined as the passage of three or more loose stool per day. On the basis of the duration of symptoms, it is further classified as acute (<14 days), persistent (14–29 days) or chronic (>30 days). Gastroenteritis, which is usually caused by a viral infection of the stomach and small bowel, is characterized by acute vomiting and diarrhoea.

Acute diarrhoea is usually caused by infections. In the West, norovirus is the commonest cause of gastroenteritis. The commonest bacterial causes, in decreasing frequency include salmonella, campylobacter, Shiga-toxin producing E. coli, vibrio and yersinia. *Clostridium difficile* infection is an important cause of diarrhoea following antibiotic or immunosuppressant treatment and in hospitalized patients. Antibiotic-associated diarrhoea occurs in the absence of *Clostridium difficile* infection and is probably due to a consequent disturbance in gut flora. The terms ‘osmotic’ (impaired fluid absorption which improves with fasting) and ‘secretory’ (excessive liquid stools >1 litre/day, which persists with fasting) are also used to classify diarrhoea with respect to the underlying physiology; examples include malabsorptive states and certain infections, respectively.

For persistent or chronic diarrhoea, the nature of the diarrhoea may give an indication as to the potential site of pathology; frequent, small-volume stools suggest a large-bowel origin, whereas less frequent but large-volume, often watery stools indicate a possible small-bowel cause:

- **Frequent, small, usually formed, stools:** colonic source such as colorectal cancer, diverticular disease, solitary rectal ulcer or normal bowel
- **Frequent, poorly formed stools:** colorectal cancer, diverticular disease, inflammatory bowel disease (Crohn’s disease and ulcerative colitis (Fig. D.9)), coeliac disease, small-bowel lymphoma, medications (laxatives, non-steroidal anti-inflammatory drugs, proton-pump inhibitors, fluoxetine), exocrine pancreatic insufficiency (cancer, chronic pancreatitis, cystic fibrosis), infiltrative disorders (scleroderma, amyloid), hyperthyroidism, Whipple’s disease

Figure D.9 Sigmoid colon with characteristic oedematous, haemorrhagic mucosa suggestive of acute ulcerative colitis.
DIARRHOEA

- Frequent, watery bowel motions: collagenous/lymphocytic colitis; bile salt malabsorption (secondary to terminal ileal resection, Crohn’s disease), infection (giardiasis, cryptosporidiosis), surgery (vagotomy, cholecystectomy, gastrectomy, small-bowel resection), enteric fistula, rapid transport (osmotic effect due to food), Crohn’s disease, arthropathy-associated terminal ileitis (ankylosing spondylitis), disaccharidase deficiency (including lactose or sucrose intolerance), post-infectious irritable bowel syndrome
- Frequent, large-volume, watery bowel motions: neuroendocrine tumours (vipoma, gastrinoma), small-bowel diseases such as enteric fistula, infections (cryptosporidiosis), drugs, surgery
- Fatty stools, which are difficult to flush away (steatorrhoea), indicate pancreatic insufficiency
- Pale, large-volume, offensive stools with bloating and borborygmia and weight loss: small bowel malabsorption (Crohn’s disease)
- Stool colour is rarely informative apart from the presence of blood, which indicates a colonic source (colitis, colorectal cancer, diverticular disease), and pale chalky-coloured stool, which indicates biliary obstruction.

As there are myriad causes for diarrhoea, it is important to relate this symptom to the patient’s history:
- Elderly (cancer, polyps, diverticular disease, ischaemic colitis)
- Middle-aged or elderly female (collagenous/lymphocytic colitis)
- Young/thin female (laxative abuse)
- Predominant constipation (spurious or overflow diarrhoea)
- Travel/local conditions (protozoal infections (giardia, entamoeba, cyclospora, cryptosporidia, strongyloides), post-infectious irritative bowel syndrome)
- Previous episodes (inflammatory bowel disease, irritable bowel syndrome)
- Medications (numerous, including antibiotics, non-steroidal anti-inflammatory drugs, proton-pump inhibitors, meftomin, colchicine, ACE-inhibitors, levodopa and fluoxetine, artificial sweeteners (sorbitol)
- Other conditions (cystic fibrosis, scleroderma)
- Recent-onset diabetes mellitus (pancreatic cancer with pancreatic insufficiency)
- Systemic features (inflammatory bowel disease, ankylosing spondylitis)
- Previous surgery (vagotomy, Polya gastrectomy, cholecystectomy, terminal ileal resection, blind loop, recent coeliac axis neurolysis, transplant)
- Family history (inflammatory bowel disease, coeliac disease, colorectal cancer, multiple endocrine neoplasia)
- Coeliac disease with recent deterioration (lymphoma)
- Symptoms associated with immunodeficiency states (HIV disease, IgA deficiency, myeloma)
- Prominent abdominal pain (structuring Crohn’s disease, ileal tuberculosis, colorectal cancer including tenesmus with rectal cancer)
- Bloating (dietary intolerances of fermentable carbohydrates)

History and physical examination can only narrow the range of likely conditions, and specific tests are always required in order to make the diagnosis. Acute diarrhoea is usually self-limiting and usually does not need investigating. Indications for investigating include severe diarrhoea (fever >38.5 °C, hypovolemia), persistent or chronic diarrhoea and risk of C. difficile infection.

Stool samples should reach the laboratory within 4 hours for parasitology (and ideally three samples) and within 12 hours for bacterial analysis. Stool can also be analyzed for elastase-1 (low levels in pancreatic insufficiency) and calprotectin (raised in any inflammatory condition and therefore useful to distinguish inflammatory from functional causes). Further evaluation with colonoscopy, including ileal and colonic biopsies, will often provide a definitive diagnosis. A flexible sigmoidoscopy is less invasive, does not require full bowel preparation and is therefore safer and can be used instead if a distal colonic cause is likely. It is of paramount importance that colorectal cancer be excluded in patients over the age of 50 years (or younger in those with a family history), even if the symptoms appear trivial.

Radiology (CT or MRI enterography) can provide additional useful information on small-bowel anatomy and transit time. Barium studies are little used nowadays. Video capsule endoscopy is rarely helpful for investigating unexplained diarrhoea, but it does provide accurate small bowel mucosal visualization in suspected or proven Crohn’s disease. (Figs D.10 and D.11).

Gastroscopy with distal duodenal biopsy will identify some enteropathies, particularly coeliac disease, but this latter condition is more easily screened for by serology (anti-endomysial or tissue trans-glutaminase antibodies). False-negative serology may occur in patients with low levels of IgA, so serum protein electrophoresis can be a helpful adjunctive test.
Bile acid malabsorption (due to terminal ileal disease or resection) is detected by the selenium homocholic acid taurine (SeHCAT) test.

Other tests for pancreatic exocrine function include 5-day faecal fat, absorptive tests (pancreolauryl test), and magnetic resonance imaging with secretin provocation or endoscopic ultrasound. Today, endoscopic retrograde cholangiopancreatography (ERCP) is less commonly used for diagnostic studies on account of the risk of inducing acute pancreatitis (about 5 per cent).

**DIPLOPIA**

Reginald Daniel

Diplopia, or double vision, is the earliest symptom apparent to the patient with dysconjugate gaze of the two eyes. It is seen with ocular paresis or following loss of normal binocular control. It is usually binocular in nature, and is rarely described in monocular form. Usually, an object is seen singly with each eye separately, but appears double when both eyes are open.

**BINOCULAR DIPLOPIA**

**Physiological diplopia**

This is a normal phenomenon in all binocular vision. Since the two eyes observe an object from different directions, the retinal images differ, but the double vision is not apparent since the dissimilar images are combined by the visual centres in the brain to form a single solid conception of the object viewed. This physiological appearance of the two retinal images is called ‘disparateness’. Such physiological diplopia is sometimes noticed by an individual who realizes that, when fixing on a particular object, other objects that are closer or further away than the object of fixation may be seen double. Patients who are anxious or unduly introspective may need reassurance about this normal phenomenon. If the centre in the brain that controls the fusion of the two images is disturbed, as after the excessive consumption of alcohol, the normal balance of the muscular mechanisms of the eyes may be lost and diplopia experienced.

**Pathological diplopia**

In normal binocular vision, both eyes are aligned so that the image of the object fixated falls upon the central and most sensitive part of the retina. Other objects form images upon more peripheral areas of the retina, and are less well observed. When an individual is looking at a particular point ahead, the image of any object lying to the right of the eyes will fall upon the nasal side of the right retina and the temporal side of the left retina, and these different areas of the retinas will always correspond and, in normal circumstances, will always be stimulated simultaneously. If the relative position of the two eyes is upset, the image of an object no longer falls upon
the two corresponding areas of the retina, erroneous forms of projection occur, and there is consequent diplopia. An examination of this diplopia leads to the ascertainment of the type of displacement of the eye. In a paralytic strabismus where one of the eyes is not able to move normally in one direction, the image seen by the affected eye, lying away from the macula of the retina, is usually less distinct. A more reliable way of distinguishing which image is the false one is by finding the direction of gaze in which the images are most displaced, and then performing a ‘cover test’ in which each eye is alternately covered; the eye that subdues the more distal image is the one that has the paralysed muscle. Alternatively, the use of a red glass test or a Maddox rod test – the former to create red and white images with white light, and the latter to create a white pinpoint image and a red line image – may be used to identify the images arising from the different eyes. For example, if the greatest horizontal separation of images is on looking to the right, either the right lateral rectus or the left medial rectus is weak. The image that is projected further from the centre is that arising in the paretic eye, and by covering one eye after the other this may be determined.

Binocular diplopia may be caused by paralysis of any extraocular muscle, and it will also be seen when there is displacement of one of the globes, as with an intraorbital tumour, abscess, haemorrhage or cavernous sinus thrombosis. It may also be seen following an operation on the extraocular muscles if the excursion of one eye is limited by scarring. Binocular diplopia arising from disease within the orbit will frequently be associated with pain or discomfort, and often with proptosis. The investigation of such syndromes is optimally undertaken with computed tomography (CT) or magnetic resonance imaging (MRI) scanning of the orbits.

Double vision may be seen in the elderly after local cerebral ischaemic events; it is usually thought to be of intrinsic brainstem origin, and is often temporary.

Patients with true paralytic strabismus will be identified as having lesions affecting the IIIrd, IVth or VIth cranial nerve, either singly or in combination, and investigations of the relevant areas may then be undertaken with CT or MRI scans, cerebrospinal fluid examination and possibly angiography. Rarely, orbital myositis will cause problems in eye movement and result in diplopia. Occasionally, neuromuscular junction disorders will affect the eye, as in ocular myasthenia. The diagnosis may be confirmed by the use of intravenous edrophonium and by single-fibre electromyography studies (Box D.7).

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**Box D.7 Differential diagnosis of binocular diplopia**

<table>
<thead>
<tr>
<th>True ocular palsy: due to lesion of IIIrd, IVth or VIth cranial nerve, or of extraocular muscles</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lesion of IIIrd, IVth or VIth nerve</td>
</tr>
<tr>
<td>• Congenital, e.g. agenesis, malformation</td>
</tr>
<tr>
<td>• Inflammation, e.g. encephalitis, neurophilis, disseminated sclerosis, herpes zoster, Tolosa–Hunt syndrome, pseudotumour of orbit</td>
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<tr>
<td>• Neoplasms, e.g. primary or secondary intracranial, meningeal tumour</td>
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<tr>
<td>• Toxic, e.g. alcohol, carbon monoxide, lead</td>
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<tr>
<td>• Vascular, e.g. thrombosis, haemorrhage, embolism, aneurysm (intracavernous), cavernous sinus thrombosis, giant-cell arteritis, diabetes, hypertension, ophthalmoplegic migraine</td>
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<tr>
<td>• Traumatic</td>
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<tr>
<td>• Raised intracranial pressure</td>
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<table>
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<tr>
<th>Lesion of extraocular muscle</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Congenital, e.g. agenesis, malformation, Möbius’ syndrome</td>
</tr>
<tr>
<td>• Traumatic, including postoperative scarring or surgical overcorrection</td>
</tr>
<tr>
<td>• Myopathy, e.g. endocrine, ocular myopathy, carcinomatous, iatrogenic</td>
</tr>
<tr>
<td>• Inflammation, e.g. ocular myositis</td>
</tr>
<tr>
<td>• Disorder of neuromuscular transmission, e.g. myasthenia gravis</td>
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</tbody>
</table>

**Relative ocular palsy: due to lesion in orbit resulting in deviation of visual axis**

| Traumatic, e.g. damage to supports of eyeball, fracture of orbital floor |
| Space-occupying lesion of orbit causing exophthalmos, e.g. tumour, haemorrhage, inflammatory lesion, mucocele, endocrine conditions |

Diplopia may occasionally be seen in patients with pituitary exophthalmos or with thyroid disturbances resulting in dysthyroid eye disease.

Ocular palsies may be central, that is due to a lesion of the nucleus or of the parenchymal portion of the cranial nerve. Alternatively, they may be peripheral, arising from lesions along the tract of the cranial nerve within the posterior or middle fossa or in the orbit itself. Lesions that can damage individual cranial nerves subserving eye movement include posterior communicating artery aneurysm, classically causing IIIrd nerve palsy, aneurysms within the cavernous sinus, raised intracranial pressure, resulting in VIth nerve palsy, and vascular lesions of the trunk of the nerve; these are the most common cause of IVth nerve palsy. The VIth nerve may also be affected near the apex of the petrous bone, when involvement with a local infective process or a nasopharyngeal tumour has to be included in the differential diagnosis.

The rare occurrence in children of episodes of ocular palsy in conjunction with unilateral headache may be recognized as ophthalmoplegic migraine. There is a rare condition in which an inflammatory or granulomatous process in the anterior portion
of the cavernous sinus or superior orbital fissure may involve any one of the nerves responsible for eye movement (Tolosa–Hunt syndrome).

The acute development of a bilateral ophthalmoplegia is most commonly seen in brainstem lesions or in post-infective cranial neuropathies (Guillain–Barré syndrome), whereas the chronic development of a bilateral ophthalmoplegia is most often seen with ocular myopathy, as in one of the mitochondrial cytopathies.

There are rare examples of pseudoparalysis of ocular muscles. In dysthyroid eye disease, a tight inferior or superior rectus muscle may limit upward and downward gaze respectively, and occasionally muscle enlargement may be demonstrated by CT scanning. The Duane syndrome, due to congenital fibrosis of the lateral rectus, causes retraction of the globe on adduction, giving rise to diplopia. However, since it is life-long, it is rarely a symptom to the patient, and is more likely to be discovered by the unsuspecting examiner.

MONOCULAR DIPLOPIA
This is infrequently found, and may be due to early cataract, or irregularity of the corneal surface, for example following inflammation.

DROP ATTACKS
Mark Kinirons
The term ‘drop attack’ is applied to a fall occurring without warning, and without loss of consciousness or post-ictal confusion. The patients are usually elderly. They suddenly drop to the floor while walking or standing; the knees buckle, there is no dizziness or other preceding symptoms, and they usually fall forward, striking the knees and sometimes the nose upon the ground. Some patients will describe a loss of power in the legs, and they are unable to raise their hands to stop the face hitting the ground, implying that normal tonus may be lost in both the lower and the upper limbs. After the episode, the patient is able to rise and move immediately. Not infrequently, such falls result in lacerations to the knees and abrasions to the face. Attacks without apparent cause may occur for a few weeks and then stop. They have a benign prognosis and there is no effective therapy.

Dyspareunia may lead to vaginismus, which is involuntary spasm of pubococcygeus muscle so that penetration is difficult or impossible. It may occur after one episode of pain, but it may become regular in an attempt to prevent subsequent episodes of pain. The pelvic floor muscles become rigid, making further attempts at intercourse painful and difficult, and thus continues the cycle. These tense muscles can be palpated on vaginal examination.

SUPERFICIAL DYSPAREUNIA
Superficial dyspareunia can be classified according to the local anatomical factors.

Vulval causes
• Infective vulvitis can occur with herpes or candidal infections. Relevant swabs and antibiotics or antiviral agents are needed.
• Atrophic changes, particularly in postmenopausal women, respond to hormone replacement therapy (HRT) or topical oestrogens. If oestrogen
is contraindicated, lubrication with KY gel or Sylk™ may be of value, as may local moisturizing preparations (e.g. Replens™).

- **Bartholinitis** may be due to local infection of the Bartholin’s gland, but this can be a site for gonorrhoea. Marsupialization has been the standard treatment to drain the cyst and create a new duct for the gland, but this is now being superseded by the use of a balloon catheter to drain any fluid and create a new permanent passage.

- **Skin conditions** affecting the vulva, including lichen planus and lichen sclerosus (see also PRURITUS VULVAE, p. 529), may cause pain as a result of the development of cracks and fissures in the skin.

- **Neoplasms**, either malignant or premalignant, may cause these symptoms and will need appropriate diagnosis and treatment.

- **Vulvodynia** is a chronic vulval discomfort that is characterized by the complaint of burning, stinging, irritation or rawness. This condition can be divided into provoked, unprovoked and a mixed picture. The diagnosis is a clinical one after excluding other causes.

**Urethral causes**

These are very much anatomical problems and are not seen very often in a general gynaecological clinic:

- **Urethritis** and **cystitis** may require local swabs or a mid-stream specimen for culture and sensitivity.

- A **caruncle** should be clearly seen on inspection, usually occurs in postmenopausal women and may become inflamed and tender.

- **Diverticulum of the urethra** is an uncommon condition to present to the gynaecologist.

**Vaginal causes**

- **Vaginismus**, as previously described.

- **Atrophic vaginitis** should be treated as above for the atrophic vulva.

- **Infective vaginitis** with Candida, Trichomonas, Herpes or Gonorrhea requires routine local swabs to be undertaken and appropriate treatment instituted.

- **Anatomical problems** may come to light in the form of vaginal atresia or imperforate hymen, which may require ultrasound scanning, to see whether there are further anatomical problems within the pelvis, and an examination under anaesthesia to determine the extent of the problem.

- **Contractures** after surgery, especially for episiotomy or perineal tear repairs, can lead to narrowing of the entrance to the vagina. The introitus may be very tight and require further surgery to relieve the local tightness caused by the original repair. Diagnosis is by clinical examination.

- **Post-radiotherapy changes**, which can be prevented to a great extent by the use of vaginal dilators around the time of initial treatment.

Disproportion in size is rarely in itself of importance as the vagina is very distensile, but if there is in addition any local lesion, the pain will be accentuated. An anal fissure and thrombosed and inflamed piles can be recognized by careful examination of the anus and rectum by a finger or speculum. Arthritis of the hips or lumbar spine may cause dyspareunia, although it may not be so well localized.

**DEEP DYSPAREUNIA**

Deep dyspareunia is due to deep stretching of the involved pelvic tissues at the time of coitus. These include a fixed retroverted uterus, the uterosacral ligaments or rectovaginal septum, or pressure on enlarged ovaries. There may be no pain on penetration and no difficulty, but coitus with deep penetration gives acute pain at the time or leads to dull aching in the pelvis after intercourse. Clinically, the symptoms can be mimicked by vaginal examination. The following are usual causes:

- **Pelvic inflammatory disease**, in which the pelvic organs may be inflamed, and adhesions may fix the tissues in place. If this is an acute episode, antibiotics can be used, and it is important to ensure that the partner is also treated. If this is a chronic picture, pelvic clearance of the genital organs may be a final-stage option.

- **Endometriosis** is a common cause of deep dyspareunia, especially when the uterosacral ligaments are involved (Fig. D.12). The degree

![Figure D.12 Endometriosis of the uterosacral ligament: laparoscopic appearance.](image)
The causes of oesophageal dysphagia can be divided into luminal, mural and extraluminal pathologies. Cancer is the single most important differential diagnosis. The luminal causes of dysphagia include: adenocarcinoma (usually distal); squamous cell carcinoma (usually proximal, but may be distal); invasive lung or mediastinal tumours (primary and metastatic); Kaposis’s sarcoma; hiatus hernia (especially the rolling type); reflux oesophagitis; peptic or lye stricture; epidermolysis bullosa (usually proximal strictures); Crohn’s disease; pemphigus; infection (viral: cytomegalovirus, herpes simplex; fungal/yeast: Candida; bacterial: tuberculosis); web formation; dysphagia lusoria’ is used to describe difficulty with swallowing, particularly in children, that results from oesophageal tethering causing dysmotility. The term ‘dysphagia lusoria’ is used to describe difficulty with swallowing, particularly in children, that results from oesophageal tethering causing dysmotility. The term ‘dysphagia lusoria’ is used to describe difficulty with swallowing, particularly in children, that results from oesophageal tethering causing dysmotility. The term ‘dysphagia lusoria’ is used to describe difficulty with swallowing, particularly in children, that results from oesophageal tethering causing dysmotility. The term ‘dysphagia lusoria’ is used to describe difficulty with swallowing, particularly in children, that results from oesophageal tethering causing dysmotility. The term ‘dysphagia lusoria’ is used to describe difficulty with swallowing, particularly in children, that results from oesophageal tethering causing dysmotility. The term ‘dysphagia lusoria’ is used to describe difficulty with swallowing, particularly in children, that results from oesophageal tethering causing dysmotility. The term ‘dysphagia lusoria’ is used to describe difficulty with swallowing, particularly in children, that results from oesophageal tethering causing dysmotility. The term ‘dysphagia lusoria’ is used to describe difficulty with swallowing, particularly in children, that results from oesophageal tethering causing dysmotility. The term ‘dysphagia lusoria’ is used to describe difficulty with swallowing, particularly in children, that results from oesophageal tethering causing dysmotility. The term ‘dysphagia lusoria’ is used to describe difficulty with swallowing, particularly in children, that results from oesophageal tethering causing dysmotility.

The investigation of dysphagia is a matter of urgency on account of the potential presence of an underlying cancer. The main tool for this investigation is gastroscopy with biopsies. Dysmotility (particularly achalasia) and eosinophilic oesophagus show few, if any, abnormalities on gastroscopy. Evidence of luminal pathology or dysmotility may be garnered from a barium swallow, particularly if swallowing is induced by combining the barium with soft foods such as bread or marshmallows.
Figure D.13 Achalasia showing gross oesophageal dilatation. The filling defects in the column of barium are due to retained masses of food.

Figure D.14 Diverticulum of the pharynx (pharyngeal pouch), demonstrated on barium swallow.

Figure D.15 Carcinoma of the oesophagus, the typical ‘rat tail’ deformity on barium swallow.

Figure D.16 Barium meal of a patient with severe dysphagia due to an extensive carcinoma of the cardia.
Oesophageal high-resolution manometry with ambulatory 24-hour pH measurement will identify acid reflux or dysmotility such as achalasia. Both computed tomography scanning and endoscopic ultrasound (Fig. D.17) permit the identification of mediastinal pathology, the latter technique allowing tissue sampling.

DYSPNOEA

Alex West

Dyspnoea is the clinical word for breathlessness. Although it is the ordinary Greek word for breathlessness, it is not an everyday term for English-speaking people. Thus, the word breathlessness, rather than dyspnoea, will be used throughout this section to emphasize that we are dealing with a sensation or symptom and not a physical sign. Most people intuitively know what is meant by breathlessness but would find it hard to describe in words. A useful definition is that offered by Comroe in 1966: ‘difficult, laboured, uncomfortable breathing; it is an unpleasant type of breathing, though it is not painful in the usual sense of the word’. Most definitions of the term involve concepts of effort and awareness of a need to breathe.

Breathlessness may be a single sensation or, like pain, several related sensations. It is not clear to what extent the breathless sensation is the same in physiological circumstances (e.g. on vigorous exercise in normal subjects) and pathological circumstances, such as in respiratory disease; it is also not clear whether the breathlessness of different disease states is qualitatively the same. These issues, although intriguing, are rarely of practical importance in the clinical situation.

Conditions associated with breathlessness can be grouped into three main categories, which are not mutually exclusive. Breathlessness occurs in conditions where there is: an increased chemical or neurological drive to breathe; an increased work of breathing; or a decreased neuromuscular power. In all these situations, there is likely to be an increased drive to breathe, whether primary or secondary, and whether or not accompanied by an actual increase in ventilation. Recent experimental work on breathlessness, in both normal subjects and patients with respiratory disease, has led to a general hypothesis for the genesis of the sensation. This suggests that breathlessness occurs when a drive to breathe exists that is abnormal, either qualitatively or quantitatively, and is translated in the medulla into a descending motor command to the respiratory muscles.

The important causes of breathlessness, in terms of the three categories described above, are listed in Table D.1. In practice, many of these conditions have their effects via more than one mechanism. There are some conditions associated with increased ventilation that are only rarely associated with breathlessness.

Conditions associated with decreased neuromuscular power are all relatively rare causes of breathlessness.

When increased ventilation is voluntary, the descending path from the cortex to the respiratory anterior horn cells bypasses the medullary respiratory centre, and breathlessness is much reduced or absent. This is evidence for the origin of the sensation in the region of the respiratory centre. In many forms
of acidosis, if the respiratory apparatus is normal, breathlessness is rare despite the increased ventilation.

**Table D.1 Causes of breathlessness**

<table>
<thead>
<tr>
<th>Conditions associated with an increased chemical or neurological drive to breathe</th>
<th>Common causes</th>
<th>Uncommon causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pulmonary oedema</td>
<td></td>
<td>Acidosis</td>
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<tr>
<td>Pulmonary embolus</td>
<td></td>
<td>Pregnancy</td>
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<tr>
<td>Pneumothorax</td>
<td></td>
<td>Cyanotic congenital heart disease</td>
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<tr>
<td>Pneumonia</td>
<td></td>
<td>High altitude</td>
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<tr>
<td>Lobar collapse</td>
<td></td>
<td>Arteriovenous fistula</td>
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<tr>
<td>Pulmonary fibrosis</td>
<td></td>
<td></td>
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<tr>
<td>Anaemia</td>
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<td></td>
</tr>
</tbody>
</table>

| Conditions associated with an increased work of breathing |
|---|---|
| **Obstructive ventilatory defects** | Common causes | Uncommon causes |
| Chronic obstructive airways disease | | Upper airways obstruction |
| Emphysema | | Byssinosis |
| Asthma | | |
| Bronchiectasis | | |
| Cystic fibrosis | | |

<table>
<thead>
<tr>
<th><strong>Restrictive ventilatory defects</strong></th>
<th>Common causes</th>
<th>Uncommon causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sarcoidosis</td>
<td></td>
<td>Large tumours</td>
</tr>
<tr>
<td>Idiopathic pulmonary fibrosis</td>
<td></td>
<td>Large hiatus hernia</td>
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<tr>
<td>Extrinsic allergic alveolitis</td>
<td></td>
<td>Lymphangitis carcinomatosa</td>
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<tr>
<td>Pneumocystosis</td>
<td></td>
<td>Connective tissue diseases leading to non-specific interstitial pneumonia</td>
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<tr>
<td>Large pleural effusion</td>
<td></td>
<td>Aspiration pneumonitis</td>
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<tr>
<td>Extensive lung resection</td>
<td></td>
<td>Infections</td>
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<tr>
<td>Chest wall deformity</td>
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<td>Asbestosis</td>
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<tr>
<td>Pulmonary oedema</td>
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<td></td>
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<tr>
<td>Left ventricular dysfunction</td>
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<table>
<thead>
<tr>
<th>Conditions associated with decreased neuromuscular power</th>
<th>Common causes</th>
<th>Uncommon causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Myasthenia gravis</td>
<td></td>
<td>Poliomyelitis</td>
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<tr>
<td>Polyneuritis</td>
<td></td>
<td>Motor neurone disease</td>
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<tr>
<td>Muscular dystrophies</td>
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**RESPIRATORY FUNCTION TESTS**

In most respiratory conditions severe enough to cause breathlessness, sophisticated tests of function are not necessary as the problem is obvious. However, respiratory function tests can be of use in determining patterns of impairment that can help in diagnosing the cause of the problem, and are particularly useful for monitoring changes in function over time or in response to treatment. Most respiratory function testing is done at rest, but important information about the cardiorespiratory system can also be obtained from measurements made during exercise.

The basic and most useful indices of lung function can be measured using a spirometer to record expired volume over time. Many machines are now available, with results produced either as flow–volume curves (Fig. D.18) or as volume–time curves (Fig. D.19). The major variables obtained from spirometry during forced expiration are the forced expiratory volume in 1 second (FEV₁), the forced vital capacity (FVC) and the peak expiratory flow rate (PEFR). Other variables that can be obtained from the same manoeuvre include flows at different points in the expiration such as at 50 per cent and 75 per cent of the vital capacity (VC). The relaxed VC
can be measured using the same equipment but with a slow expiration. The PEFR can also be measured by simpler, cheaper peak flow meters; these are convenient when repeated estimates of PEFR are required, for example to diagnose variable airflow limitation in asthma. To estimate other lung volumes, it is necessary to use helium dilution to give an estimate of residual volume (RV, the volume remaining in the lungs at the end of a full expiration). Total lung capacity (TLC) is the sum of the RV and the VC.

Another aspect of lung function is gas exchange between the alveoli and the blood. This can be assessed by measuring the transfer factor for carbon monoxide. Of course, arterial oxygen tension is also a good indication of gas exchange, and a fall during exercise is a sensitive means of detecting interstitial lung disease affecting gas exchange. Measurement of arterial blood gas tensions requires arterial puncture, which can be uncomfortable and even hazardous and, in the absence of an indwelling arterial line, difficult to repeat frequently.

It is easy to measure arterial oxygen saturation non-invasively using oximeters. These function by shining different wavelengths of light through the tissue (usually the ear lobe or fingertip); they can be left in place for several hours, for example during exercise testing or during sleep. Similarly, it is possible to measure arterial blood carbon dioxide tension (PCO₂) non-invasively, using a sensor strapped to the skin.

Cardio-pulmonary exercise testing can assess the functioning of the cardio-respiratory system under stress. Commonly, exercise of progressively greater severity is performed on the treadmill or bicycle ergometer. Variables such as ventilation, heart rate, oxygen consumption and carbon dioxide production can be measured; it is also possible to obtain the subject’s assessments of degree of breathlessness during the test. This can sometimes elucidate the cause of breathlessness on exercise in someone with relatively normal results at rest (e.g. an exercise-induced cardiac arrhythmia or a profound fall in arterial oxygen tension, PO₂, due to a reversal of an arteriovenous shunt).

The pattern of lung function abnormality is particularly useful in deciding what type of disease is present. In airways obstruction, such as is found in chronic obstructive pulmonary disease (COPD) or asthma, spirometry typically shows a reduction in FEV₁, and in the FEV₁/FVC ratio. If a flow–volume curve is examined, the descending part of the curve tends to be concave due to reductions in flow at low lung volumes (see Fig. D.18). In some cases, there is quite a marked reduction in FVC, due to air-trapping during the forced expiration. This is especially likely in COPD. Spirometry alone in such
cases cannot distinguish between obstruction with air-trapping and a mixed obstructive–restrictive defect. Restrictive lung conditions, such as diffuse pulmonary fibrosis, typically produce a reduction in FVC but have less effect on the FEV₁ so that the FEV₁/FVC ratio tends to be high. A typical flow–volume curve in restrictive lung disease is shown in Figure D.18.

Additional tests are not always necessary to distinguish between obstructive and restrictive conditions. Static lung volumes are uniformly reduced in restrictive conditions, with a normal RV/TLC ratio, and there is frequently a reduction in gas transfer factor. In obstructive conditions, it is common to find that the relaxed VC is greater than the FVC, and that the RV/TLC ratio is increased, both due to air-trapping. If emphysema is present, destruction of lung tissue often results in a reduction in gas transfer factor.

Blood gases can become abnormal in any severe lung disease. In respiratory failure secondary to chronic obstructive airways disease, the PO₂ is reduced and there is a tendency for the PCO₂ to increase due to alveolar hypoventilation. In asthma, a reduction in PO₂ is the main feature, and the PCO₂ tends to be low due to hyperventilation in response to the hypoxaemia, only rising as a late event in life-threatening asthma. In restrictive lung conditions, a low PO₂ with a low PCO₂ is the usual pattern. In all these situations, including restrictive lung conditions, the hypoxaemia is mainly secondary to a mismatch between the ventilation and perfusion of the alveoli rather than to an actual diffusion defect.

The various conditions associated with breathlessness will be discussed under the headings used in Table D.1. Many of these produce breathlessness by several different mechanisms, but they are classified according to the primary mechanism operating in most cases.

**CONDITIONS ASSOCIATED WITH AN INCREASED CHEMICAL OR NEUROLOGICAL DRIVE TO BREATHE**

A characteristic of increased ventilation due to chemical or neurological stimulation is a high respiratory frequency and a low arterial PCO₂. Hypoxia and acidosis are the important chemical drives to breathing. Potential sources of neurological drives to breathing include pulmonary receptors, chest wall receptors and other skeletal muscle receptors. Neurological drives to breathing are probably important in parenchymal lung conditions associated with breathlessness, such as pulmonary oedema and pulmonary infarction secondary to pulmonary embolus. There may also be a mechanism for the breathlessness associated with pneumonia and lobar collapse (e.g. secondary to bronchial obstruction produced by bronchogenic carcinoma). It has been demonstrated that the increased ventilation associated with lobar collapse occurs even when the collapsed segment is small and there is little or no hypoxia. Breathlessness, due perhaps to stimulation of lung receptors, is a prominent early symptom of *Pneumocystis jiroveci* (formerly known as *Pneumocystis carinii* pneumonias) associated with HIV infection (Fig. D.20). This condition typically has an insidious onset and a long history stretching back weeks or months with a dry cough a prominent feature; the chest radiograph may be normal, but arterial oxygen desaturation on exercise is usual. The breathlessness that accompanies a relatively small spontaneous pneumothorax is often quite marked, especially in the early stages, and cannot be explained by mechanical difficulties with ventilation. Again, it is possible that stimulation of lung receptors plays a part.

Hypoxia from any cause can lead to breathlessness via stimulation of ventilation, for example in cyanotic congenital heart disease, high altitude (mountaineers) and a tendency for the PCO₂ to increase due to alveolar parenchymal lung disease such as the various types of pulmonary fibrosis. It is not clear whether hypoxia can

![Figure D.20](extensive bilateral shadowing due to *Pneumocystis jiroveci* infection.)
DYSPNOEA

act as a direct stimulus to breathlessness, independent of its effects on ventilation; experiments have produced conflicting results.

In cyanotic congenital heart disease, such as Fallot’s tetralogy, the mixed venous oxygen saturation falls on exercise due to increased extraction by the exercising muscles (this being a normal phenomenon); some of this venous blood passes into the aorta via the ventricular septal defect, causing a steep fall in arterial oxygen saturation. This stimulates ventilation and leads to severe breathlessness on exercise. Children with Fallot’s tetralogy are often observed to squat. This produces some obstruction of venous return from the exercising muscles of the legs, reducing the fall in arterial oxygen saturation and the resultant increased ventilation and breathlessness.

Severe arterial hypoxaemia occurs in respiratory distress syndrome of the newborn. Due to a lack of pulmonary surfactant, parts of the lung fail to expand, and there is marked mismatching of ventilation and perfusion in the lungs. Perfusion of unventilated segments contributes to the arterial hypoxia. There is profound stimulation of ventilation, with respiratory rates of up to 100 per minute. It is, of course, only speculation to suggest that this is accompanied by breathlessness in these infants.

Hypoxia is a common feature of many types of parenchymal lung disease; it is more often due to mismatching of ventilation and perfusion than to an actual block to diffusion (see above). A more important cause of breathlessness in many of these conditions is the increased work of breathing, due to decreased compliance of the lung tissue, and these points are discussed more fully below.

Anaemia is frequently accompanied by a complaint of breathlessness. The mechanism is probably inefficient oxygen delivery to exercising muscles, with a resultant increase in anaerobic metabolism and lactate production, which drives ventilation. The breathlessness that sometimes accompanies pregnancy and arteriovenous fistula may in part be due to a similar mechanism, with changes in distribution of the cardiac output leaving some muscle groups relatively hypoxic. In late pregnancy, mechanical difficulties with ventilation are likely to be more important. Other causes of acidosis also produce an increase in ventilation, such as renal failure and diabetic keto-acidosis. In practice, these conditions are rarely accompanied by breathlessness unless there is concomitant lung disease. In the case of keto-acidosis, this may be because consciousness is obtunded, but it remains unclear why there is relatively little breathlessness associated with other causes of acidosis.

Isocapnic voluntary hyperventilation is not accompanied by breathlessness in the experimental situation, as discussed above. Similarly, breathlessness is a relatively rare complaint in the hyperventilation syndrome; symptoms such as nausea, light-headedness and paraesthesiae are much more common. Although it has been suggested that some people with lung disease have disproportionate dyspnoea, associated with various psychiatric symptoms, true psychogenic dyspnoea is a very rare phenomenon and should never be diagnosed without very thorough investigation, including exercise testing. It should be noted, however, that the sighing respiration of cardiac neurosis may be referred to by the patient as ‘breathlessness’.

CONDITIONS ASSOCIATED WITH AN INCREASED WORK OF BREATHING

Obstructive ventilatory defects

The term chronic obstructive pulmonary disease refers to the situation where there is chronic, more or less irreversible, obstruction of the airways, especially the smaller airways. It is sometimes referred to by other names, such as chronic airflow limitation, chronic obstructive airs disease, chronic bronchitis and emphysema or chronic obstructive lung disease. The most important association is with cigarette smoking, but general atmospheric pollution also plays a part, and it is common in some dusty occupations. The term ‘chronic bronchitis’ should probably be used in relation to the patient’s history where it refers to chronic mucus hypersecretion, so a daily productive cough for more than 3 months of the year for 2 consecutive years.

COPD remains a very common condition and an important cause of breathlessness due to the increased drive to breathe required to overcome the increased resistance to airflow. It is diagnosed by the finding of airflow limitation, not reversible by bronchodilators, usually with a history of cigarette smoking and productive cough. The main differential diagnosis is asthma (see below).

In advanced COPD, the chest becomes chronically overinflated with an increased anteroposterior diameter – the so-called ‘barrel chest’. The chest radiograph shows a loss of normal lung markings, flattened domes of the diaphragm and sometimes the presence of bullae. The lung function defect is obstructive, with collapse of the airways during expiration partly due to a lack of support from the surrounding lung tissue. Other common lung function
findings are an increased RV, an increased TLC, an increased RV/TLC ratio and a decreased transfer factor. Occasionally, large bullae may act as space-occupying lesions and add a restrictive component to the picture. There is debate about whether resection of the bullae relieves breathlessness in these cases. Emphysema is defined as an increase in size, due to destruction of the alveolar walls, of the air spaces distal to the terminal bronchioles (Fig. D.21). It is a pathological diagnosis (but can also be clearly seen on a CT scan of the chest) and is part of the pathology of COPD. A particular type of emphysema is that related to inherited deficiency of alpha-proteinase inhibitor; the emphysema is panacinar rather than centrilobular, and it tends to affect the upper, rather than the lower, lobes.

Breathlessness due to chronic irreversible obstructive airways disease presents as a major feature in a well-known clinical picture. Expiratory wheezes at the mouth and on auscultation are common but not invariable. Respiratory failure, with hypoxia and often hypercapnia, develops. Cor pulmonale, with peripheral oedema, develops in some cases. Two extremes have been described, depending on the degree to which the respiratory centre retains its sensitivity to carbon dioxide. Where sensitivity is lost, the PCO$_2$ rises, oedema, and there is cyanosis due to hypoxia and sometimes secondary polycythaemia; this is the so-called ‘blue bloater’. Where carbon dioxide sensitivity is retained, there tends to be a greater drive to breathe, with correspondingly more severe breathlessness but relatively normal blood gases until a late stage of disease; this is described as the ‘pink puffer’. In practice, there is considerable overlap between these two extremes.

Bronchiectasis is associated with breathlessness if there is appreciable airways obstruction present and if the lung destruction is extensive. The diagnosis is suggested by a history of production of copious purulent sputum, often with haemoptysis. Clubbing may be present with extensive disease. CT scanning of the chest will confirm a diagnosis if not clearly seen on the chest X-ray (CXR). The lung disease of cystic fibrosis in children is similar to bronchiectasis of adults, with destruction of lung tissue and airways obstruction. One common cause of breathlessness in all age groups is the episodic airways obstruction of asthma. The cardinal feature of asthma is the variable nature of the airways obstruction, with partial or complete reversal after bronchodilator treatment. The typical history is of episodes of breathlessness, accompanied by wheezing, coughing or chest tightness, occurring especially at night and after exercise. During an attack, there is overinflation of the chest, wheezing and rapid ventilation. The breathlessness is made worse by the ventilatory stimulation from the accompanying hypoxia. A palpable ‘paradox’ of the arterial pulse is often present during a severe episode due to the large intrathoracic pressure swings. There are some particular causes of asthma that merit mention.

Bronchopulmonary aspergillosis may provoke episodes of asthma or worsen pre-existing asthma. The chest radiograph may show transient pulmonary infiltrations, and there is often an eosinophilia in the peripheral blood. Occupational asthma has been recognized with increasing frequency and should always be considered, especially when asthma fails to respond to usual treatments. A careful occupational history should be taken in all cases of asthma.

Airways obstruction is the main feature of byssinosis, a lung condition caused by the inhalation of cotton and other vegetable dusts. The typical history is of breathlessness and chest tightness occurring on the first day back at work after a break, and then gradually progressing so that it is constantly present. The late stages of byssinosis are indistinguishable from other causes of chronic obstructive airways disease.

Upper airways obstruction is an uncommon cause of breathlessness, but it is important to recognize because it can be rapidly fatal, although quickly relieved by intubation or tracheostomy. Obstruction at the level of the larynx may be due to severe laryngitis, which may still be secondary to diphtheria, or to laryngeal oedema as part of an allergic reaction. Severe laryngotracheobronchitis, as occurs in croup,

Figure D.21 Chest radiograph showing advanced emphysema with hyperinflated lungs and a flattened hemidiaphragm.
can rapidly produce life-threatening respiratory obstruction and breathlessness, and respiratory embarrassment can occur with severe **tonsillitis**. Obstruction can also be due to an impacted **foreign body** or to problems with the vocal cords themselves, as in **bilateral abductor paralysis**. Obstruction at the level of the trachea can be produced by intrinsic **carcinoma of the trachea** or by external compression from **carcinoma of the thyroid**, **haemorrhage into a thyroid cyst**, **neoplastic glands in the neck or mediastinum**, **aortic aneurysm** or **dermoid tumour** of the mediastinum. Obstruction of the upper airways may be accompanied by inspiratory stridor, and the flow–volume loop shows a characteristic truncation of the inspiratory part of the loop.

**Restrictive ventilatory defects**

The most obvious form of restrictive ventilatory defect is when there has been actual loss of lung tissue, following **surgical resection** of a lung or lobe. Breathlessness following pneumonectomy is more likely when there is disease of the remaining lung, such as obstructive lung disease in a smoker who has had a pneumonectomy for lung cancer. Space-occupying lesions can produce a similar effect. Examples are a **large pleural effusion** or **spontaneous pneumothorax**. Part of the breathlessness associated with these conditions is probably due to a neurological drive to breathe, but the effusions also produce a restriction of lung tissue when large. A pleural effusion may be large before it produces noticeable breathlessness in young, healthy adults, but even a small effusion can lead to a worsening of breathlessness when it occurs in association with left ventricular failure, when there may already be pulmonary hypertension and oedema. Common features of a spontaneous pneumothorax are a sharp chest pain and moderate breathlessness. A tension pneumothorax, actively increasing in size and compressing lung tissue, produces extreme breathlessness and often cardiovascular collapse and requires urgent relief through insertion of a cannula. Physical signs of a large pneumothorax are reduced movement and reduced (or absent) breath sounds on the affected side. Large **tumours** can behave as space-occupying lesions, a particularly unpleasant example being a **mesothelioma** secondary to asbestos exposure. At post-mortem examination, the tumour is often seen to have replaced most of the original lung volume. A large **hiatus hernia** can occasionally cause breathlessness by acting as a space-occupying lesion within the chest.

Lung restriction can result from conditions of the chest wall. Deformity of the thoracic cage, such as occurs in severe **kyphoscoliosis**, **ankylosing spondylitis** or after a **thoracoplasty**, reduces the inspiratory volume that can be achieved, and may be associated with breathlessness as a result. Extensive pleural fibrosis or calcification can similarly produce a restrictive ventilatory defect; this may be found following **tuberculosis** or secondary to **asbestos exposure**. In severe cases, the lungs become encased in a rigid shell of pleura, sometimes called a ‘cuirass’. Restrictive ventilatory defects due to conditions outside the lung itself are characterized by reduced lung volumes, but a normal or even increased transfer factor for carbon monoxide. This is because the lung tissue itself is normal, albeit compressed, and there may be a higher blood flow in the smaller volume of tissue, causing the increase in transfer factor.

The largest group of conditions producing restrictive ventilatory defects is those involving the lung parenchyma itself. **Idiopathic pulmonary fibrosis** (usual **interstitial pneumonia** – UIP) is a generalized condition of the lung parenchyma that begins as an inflammatory process and frequently progresses to established fibrosis. There are a number of interstitial lung diseases, differing in their rate of progression and response to treatment with corticosteroids or immunosuppressants. The most acute form, originally known as the Hanman–Rich syndrome but now known as **acute interstitial pneumonia** (AIP), progresses to severe disability and death within weeks or months. Other forms can be present for years before causing severe symptoms. Common features are breathlessness on exertion, dry cough, clubbing of the fingers (and toes) and inspiratory crackles, especially at the lung bases. The chest radiograph shows reticulonodular shadowing, usually more marked in the lower zones.

In **idiopathic pulmonary fibrosis**, as the name suggests, the cause is unknown. A somewhat similar clinical picture can develop in some cases of extrinsic allergic alveolitis (or hypersensitivity pneumonitis (HP)), which develops in response to inhaled organic dusts. The major mechanism is a type III allergy, with IgG antibodies against the allergen present in the blood. The best-known example of typical extrinsic allergic alveolitis (EAA) is farmer’s lung. Acute episodes follow heavy exposure to the allergen, in this case spores of fungi such as **Micropolyspora faeni** contaminating damp hay. There is fever, breathlessness, cough and sometimes cyanosis, and crackles are audible in the chest. Hypoxia and a restrictive ventilatory defect are present, and there are scattered small, nodular shadows on the chest radiograph. Such acute episodes are often difficult to distinguish from atypical
pneumonia, although the presence of precipitating antibodies can help. If exposure to the responsible agent continues, chronic disease develops. This is characterized by persistent breathlessness and dry cough, sometimes with clubbing and cyanosis. Airways obstruction can develop in addition to the restrictive defect. The diagnosis is helped if there is a history of preceding acute episodes. In some types of EAA, these are rare; for example, in budgerigar fancier's lung, an insidious development of fibrosis is common, probably because of the continuous low level exposure in this case. There are numerous other causes of EAA, including: fungal spores, involved in the bagassosis of sugar-cane workers, in suberosis of cork workers, in sequoiosis of redwood sawyers, and in lung disease of mushroom workers, malt workers, maple-bark strippers and cheese makers; animal proteins in pigeon fancier's lung; and bacterial or amoebal proteins in humidifier fever (as distinct from legionnaires' disease, where lung infection is involved).

Extrinsic allergic alveolitis has some clinical and pathological features in common with sarcoidosis, although in the latter no causative agent has been identified. Acute, subacute and chronic forms of sarcoidosis are described. The acute form may have little in the way of respiratory symptoms, and is characterized by bilateral hilar lymphadenopathy, often with fever, erythema nodosum and arthralgia. In the subacute and chronic forms, there is a granulomatous infiltration in the lungs with increasing breathlessness. The chest radiograph shows pulmonary infiltration, said to be characteristically perihilar but often more generalized (Fig. D.22). Sarcoidosis is a systemic condition, and many organs apart from the lungs can be affected, including the eyes (iritocyclitis), the lacrimal and salivary glands, the lymph nodes, the liver, the skin and the central and peripheral nervous systems. A useful diagnostic feature of sarcoidosis, and EAA, is the presence of a large number of lymphocytes in the bronchoalveolar lavage fluid but more usefully a bronchial or transbronchial biopsy may reveal non-caseating granuloma which is characteristic of sarcoid. In idiopathic pulmonary fibrosis, on the other hand, the fluid contains increased numbers of neutrophils.

Pneumoconiosis is a general term for lung disease due to dust inhalation, but it is used mainly to describe the conditions associated with inhaling inorganic dusts where allergy is not an important mechanism. Silicosis is still an important disease worldwide. Breathlessness develops gradually, and the restrictive pattern of deficit is frequently complicated by airways obstruction due to airways disease and lung distortion by fibrotic masses.

Figure D.22 (a) Chest radiograph of acute sarcoid with mediastinal and hilar lymphadenopathy (arrowed). (b) Chest radiograph of chronic sarcoid with bilateral upper and mid-zone infiltration (arrowed).

The fibrosis of silicosis is nodular rather than diffuse, and the chest radiograph shows distinct nodules that tend to coalesce to form large masses. In coal worker's pneumoconiosis, the chest radiograph shows a background of small nodular opacities, with large masses as the disease progresses (Fig. D.23). Disability is due mainly to the associated emphysema, which may be severe. Asbestosis is a very common cause of restrictive lung disease in the industrialized world (Fig. D.24). Symptoms and signs of lung disease tend
to develop after exposure over many months or years. The lung fibrosis associated with asbestos exposure is diffuse and affects especially the lower lobes. It is frequently accompanied by pleural thickening or plaques, which may calcify. An important complication is lung cancer, or more rarely mesothelioma. The restrictive lung function deficit and breathlessness of asbestosis is only poorly related to the severity of the radiographic changes. A number of other dusts can produce forms of pneumoconiosis. Berylliosis, caused by exposure to beryllium, has many features in common with sarcoidosis, and sometimes responds to treatment with corticosteroids.

Carcinoma of the lung (see Fig. C.27, p. 114) rarely causes breathlessness as an early feature, but breathlessness in more advanced disease may be due to a lobar collapse (as discussed above), to multiple metastases in the lungs, or to lymphatic involvement producing lymphangitis carcinomatosa. This may also be due to the spread of extrapulmonary tumours, and is associated with very severe breathlessness.

Most of the generalized connective tissue diseases can affect the lungs including polymyositis and mixed connective tissue disease. On CT scanning they may have a so-called NSIP (non-specific interstitial pneumonia) or an ‘organizing pneumonia’ type pattern. Systemic lupus erythematosus can involve the pleura, with recurrent episodes of pleurisy, but it can also lead to lung fibrosis. Systemic sclerosis (scleroderma) produces a diffuse interstitial pulmonary fibrosis, which can go on to produce cor pulmonale. A further cause of lung damage in this condition is recurrent aspiration if the oesophagus is also affected. Pulmonary involvement in rheumatoid disease can be seen as diffuse fibrosis or rheumatoid nodules in the lung can occasionally occur. Caplan’s syndrome is a type of coal worker’s pneumoconiosis that occurs in men who have the rheumatoid diathesis. Multiple rounded opacities occur on a background of scant small nodules; calcification and cavitation are quite common (Fig. D.25). The lung conditions related to connective tissue diseases can occur without overt signs of the rheumatological initially but may be detected through underlying specific blood tests sometimes.

Lung destruction as a result of infections can produce breathlessness secondary to a restrictive defect. Tuberculosis, with extensive cavitation, is often associated with breathlessness before adequate treatment; the breathlessness may persist after bacteriological cure due to the loss of lung tissue. Klebsiella pneumonia frequently destroys lung tissue,
and severe cases may be left with breathlessness after recovery from the infection.

Other conditions of the lung parenchyma that can produce a restrictive ventilatory defect include schistosomiasis, fungal infections (e.g. blastomycosis and coccidioidomycosis) and alveolar proteinosis. Poisoning with the weedkiller paraquat leads to severe breathlessness due to an obliterative bronchiolitis. One important group of conditions in which breathlessness occurs at least partially due to a restrictive ventilatory defect is that causing pulmonary oedema. Cardiac causes of pulmonary oedema include valvular lesions (e.g. mitral stenosis) and left ventricular failure. The most common cause of left ventricular failure is ischaemic heart disease; other causes are hypertensive heart disease and non-ischaemic cardiomyopathies. As the pulmonary venous and capillary pressures rise, the lungs become stiffer; the accumulation of fluid in the interstitial tissue and in the alveoli increases the lung stiffness. Thus, the work of ventilation is increased. In addition, gas transfer is impaired leading to increasingly severe arterial hypoxaemia, which further drives ventilation. Breathlessness on exertion is accompanied by orthopnoea and bouts of nocturnal breathlessness (so-called paroxysmal nocturnal dyspnoea). The breathlessness that is frequently the symptom which limits exercise in heart disease with left ventricular dysfunction is not fully understood. The mechanism just outlined operates in some cases, but other mechanisms – perhaps involving neurological drives to breathe – may also be important.

Acute pulmonary oedema, as every house officer knows, is a dramatic condition, with severe breathlessness developing rapidly, accompanied by a cough productive of frothy, pink sputum. There are widespread crackles in the chest, and wheezing is often a prominent feature due to oedema of the bronchial walls. It can sometimes be difficult to differentiate between acute asthma and pulmonary oedema; radiographic appearances and the finding of mitral stenosis or a cause of left ventricular failure can be helpful. It is always worth remembering that an elderly patient with severe breathlessness might be suffering from acute asthma rather than pulmonary oedema secondary to cardiac disease. The radiographic features of pulmonary venous hypertension include engorgement of the upper lobe veins and septal lymphatic lines. In acute pulmonary oedema, there is enlargement of the hilar shadows and fan-shaped mid-zone opacities (Fig. D.26).

Pulmonary oedema can be precipitated by the infusion of saline. This is unlikely to occur with a normal heart and adequate renal function, but it can readily occur in oliguric renal failure. Other non-cardiac causes of pulmonary oedema include cerebral vascular accidents, salicylate poisoning, hypersensitivity to radiographic contrast media, and adult respiratory distress syndrome when the pulmonary vasculature becomes abnormally ‘leaky’. Pulmonary oedema can be caused by the inhalation of smoke or other irritant gases, such as chlorine. Aspiration pneumonitis (Mendelson’s syndrome) is similar to pulmonary oedema. It is due to the aspiration of gastric acid, and the symptoms are virtually the same as for acute pulmonary oedema.

Figure D.25 Chest radiograph (a) and high-resolution computed tomography scan (b) of the lungs showing grossly dilated bronchi (arrowed) in a patient with allergic bronchopulmonary aspergillosis.
CONDITIONS ASSOCIATED WITH DECREASED NEUROMUSCULAR POWER

In these conditions, an increased drive to the functioning respiratory muscles is required in order to attempt to maintain ventilation. Weakness of the respiratory muscles may be due to primary muscle disorders or to neurological disease. Examples include myasthenia gravis, polyneuritis, poliomyelitis, motor neurone disease and muscular dystrophies. Despite the drive to breathe, hypoventilation is a feature of these conditions. It is also a feature of the respiratory failure associated with end-stage COPD, in particular. The hypoventilation associated with rare lesions of the respiratory centre itself is not usually associated with breathlessness, nor is the hypoventilation found in the obesity hypoventilation syndrome or Pickwickian syndrome.

DIFFERENTIAL DIAGNOSIS OF ACUTE BREATHLESSNESS

Acute severe breathlessness is a common medical emergency that is alarming for the patient and medical attendant alike. The urgency of the situation may be such that no time is available for radiographic or other investigation, and the initial diagnosis must be made on the basis of clinical examination. The differential diagnosis will be discussed in terms of the physical signs which may be elicited in such an examination. Simple inspection can be informative. If the patient is deeply cyanosed with engorged cervical veins, struggling for breath with violent movements of the larynx, laryngeal or upper tracheal obstruction should be considered. The stridor of upper airways obstruction is heard during both inspiration and expiration, and this should be distinguished from the expiratory wheeze of ordinary bronchial obstruction. The possible causes of upper airways obstruction are discussed above. The mucosal pallor of severe anaemia as a cause of breathlessness may also be noted on inspection. Examination of the arterial pulse may reveal uncontrolled atrial fibrillation or other arrhythmias, which can lead to sudden breathlessness, especially if there is valvular or other cardiac disease. Cardiac tamponade is associated with marked ‘paradox’ of the arterial pulse and elevation of the jugular venous pressure, with a paradoxical rise in pressure during inspiration. It may be secondary to intrapericardial haemorrhage from malignant metastases; in association with cardiac rupture from a myocardial infarction, it is usually rapidly fatal. The venous pressure is also markedly elevated following a massive pulmonary embolus, another cause of acute severe breathlessness.

Mediastinal displacement, as indicated by lateral displacement of the trachea, can be a valuable physical sign in acute breathlessness. In massive collapse of a lung, for example postoperatively, the trachea is displaced towards the affected side. The trachea is deviated away from the side of the lesion in pleural effusion and tension pneumothorax. The most likely fluid to accumulate rapidly enough in the pleural space to cause acute breathlessness is blood; haemothorax may follow chest trauma or even occur after pleural aspiration.

In acute pulmonary oedema, the venous pressure is usually also elevated, and there may be signs on examination of the heart, such as a murmur of mitral stenosis or a gallop rhythm in left ventricular failure. Auscultation of the lungs usually reveals widespread crackles in pulmonary oedema and expiratory wheezes in acute asthma; in very severe asthma, the wheeze can disappear because of gross reduction of air movement. If acute breathlessness is due to severe pulmonary infection, features such as pyrexia, purulent sputum and signs of consolidation in the lungs can be helpful diagnostically. Evidence of severe immunosuppression, such as oral candidiasis, oral hairy leukoplakia, or evidence of extreme weight loss should prompt consideration of Pneumocystis pneumonia complicating HIV infection.

DYSURIA

Ben Challacombe

Dysuria relates to either painful or difficult micturition, and there is some variation regarding this definition. In the UK, dysuria is the term applied to pain,
Dysuria during micturition may be due to acute or chronic cystitis and acute or chronic prostatitis. Acute prostatitis is usually due to bacterial infection of the prostate gland, and presents in younger men with symptoms of a febrile acute urinary infection. Common organisms include enterococci, Escherichia coli, Klebsiella pneumoniae and Proteus mirabilis, while STI-related organisms are Chlamydia, the gonococcus and Trichomonas. Increasingly, this condition is due to a recent transrectal, ultrasound-guided prostate biopsy as part of the investigation of prostate cancer. Acute prostatitis is rare in young boys but more common in men aged 20–35 with multiple sexual partners. Also at high risk are those who practise anal intercourse, especially without using condoms, men with a urinary catheter and those with benign prostatic hyperplasia.

The diagnosis is made on the history and confirmed by clinical examination, when the prostate is swollen and exquisitely tender, making the examination difficult. The prostate should never be massaged in acute prostatitis as a septic episode may result. Chronic prostatitis is an intractable condition in which pain is felt in the perineum and/or at the base of the penis, and may be felt on micturition. The prostate is tender on examination. Prostatic massage should be used to obtain fluid for culture of the prostatic fluid and microscopy for inflammatory cells. Tuberculous cystitis and infection with schistosomiasis may cause dysuria and, although rare, should be considered when there is a sterile cystitis or a history of foreign travel.

When a bladder calculus comes into contact with the trigone and bladder neck, pain may be referred to the urethra. Urethral pain may be felt when a ureteric stone is passing down the distal ureter, particularly when the stone is close to the bladder, i.e. in the intramural ureter, or when the stone is passed after a bout of ureteric colic. If a stone gets stuck in the urethra, as may happen when a ureteric or bladder stone is passed, urethral pain will occur, with the sudden onset of obstruction and then retention of urine. As the meatus is the narrowest portion of the urethra, a stone may lodge in the meatus, and may be felt in the distal urethra or felt and seen at the meatus. A bladder tumour close to the bladder neck may cause discomfort on voiding or obstruction, and carcinoma-in-situ of the bladder commonly results in prolonged dysuria and irritative lower urinary tract symptoms (Fig. D.27). A blood clot that has developed from bleeding in the bladder or upper urinary tract may also cause dysuria and discomfort when passed.

There are several causes of mild dysuria due to local inflammation or contact dermatitis, including irritation...
DYSURIA

from chemicals in soaps, bubble baths, spermicides and douches. Pain in the perineum can be caused by prostatitis. It may occur in infiltrating prostate cancer and urethritis caused by urethral infection. Meatal pain may be caused by genital warts or distal urethritis caused by sexually transmitted diseases. Pain in the foreskin may be seen in phimosis, paraphimosis or balanoposthitis/balanitis.

Pain felt in the bladder area is not true dysuria but may also be present during an acute episode of cystitis. Bladder pain can also be experienced in the presence of a bladder stone, when the stone bounces up and down on the trigone during the act of micturition. The trigone is a particularly sensitive area of the bladder and can therefore give rise to pain when involved by bladder or prostatic carcinoma and in acute or chronic prostatitis.

DYSURIA IN CHILDREN

Acute cystitis will cause severe voiding pain, and the child or infant will cry when voiding occurs. Meatal ulceration and acute balanitis, in the male infant, will also cause pain on micturition. Bear in mind the possibility of child sexual abuse.

*Figure D.27* Radical cystectomy specimen with an inked margin and ureteric stent showing a large transitional cell tumour near the trigone, which presented with dysuria, urinary frequency and urgency.
Earache (otalgia) may be caused by a number of conditions that affect the external and middle ear. Pain may also be referred to the ear as a result of disease processes at distant sites. These sites share sensory innervation from the same cranial nerves and spinal roots that supply the ear. Possible causes of earache are listed in Box E.1.

**AURICLE**

Causes in the auricle are usually visible. They include direct trauma, haematoma, furuncles, erysipelas and perichondritis. The distinction between erysipelas and perichondritis can be made by inspection of the lobule as this is not affected in perichondritis. The infectious causes respond to systemic antibiotics. Haematomas need to be drained as soon as possible to avoid permanent damage.

**Box E.1 Causes of earache**

<table>
<thead>
<tr>
<th>Local</th>
<th>Middle ear</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Auricle</td>
<td>• Acute otitis media</td>
</tr>
<tr>
<td>– Trauma</td>
<td>– Advanced chronic otitis media</td>
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<tr>
<td>− Direct</td>
<td>− Mastoiditis</td>
</tr>
<tr>
<td>− Haematoma</td>
<td>− Malignant disease</td>
</tr>
<tr>
<td>– Furunculosis</td>
<td>− Otitis media with effusion (rarely)</td>
</tr>
<tr>
<td>– Sebaceous cyst</td>
<td>− Mastoiditis</td>
</tr>
<tr>
<td>– Perichondritis</td>
<td>− Malignant disease</td>
</tr>
<tr>
<td>– Erysipelas</td>
<td>− Otitis media with effusion (rarely)</td>
</tr>
<tr>
<td>– Chondrodermatitis nodularis chronicis helices is a benign nodular lesion of unknown aetiology that affects the lateral edge of the pinna (Fig. E.1). It is extremely painful and sometimes needs to be excised. Malignant disease and squamous or basal cell carcinoma are frequently seen in the elderly and, when advanced, become painful (Fig. E.2). Most can be successfully treated by either surgery or radiotherapy or a combination of the two.</td>
<td></td>
</tr>
<tr>
<td>− Malignant disease</td>
<td>− Otitis media with effusion (rarely)</td>
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<tr>
<td>− External auditory meatus</td>
<td>− Mastoiditis</td>
</tr>
<tr>
<td>− Diffuse otitis externa</td>
<td>− Malignant disease</td>
</tr>
<tr>
<td>− Infective</td>
<td>− Otitis media with effusion (rarely)</td>
</tr>
<tr>
<td>− Bacterial</td>
<td>− Mastoiditis</td>
</tr>
<tr>
<td>− Fungal</td>
<td>− Malignant disease</td>
</tr>
<tr>
<td>− Viral</td>
<td>− Otitis media with effusion (rarely)</td>
</tr>
<tr>
<td>− Reactive</td>
<td>− Mastoiditis</td>
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<tr>
<td>− Eczematous</td>
<td>− Malignant disease</td>
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<tr>
<td>− Seborrhoeic dermatitis</td>
<td>− Otitis media with effusion (rarely)</td>
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<tr>
<td>− Neurodermatitis</td>
<td>− Mastoiditis</td>
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<tr>
<td>− Malignant otitis externa</td>
<td>− Malignant disease</td>
</tr>
<tr>
<td>− Benign necrotizing osteitis</td>
<td>− Otitis media with effusion (rarely)</td>
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<tr>
<td>− Wax impaction</td>
<td>− Mastoiditis</td>
</tr>
<tr>
<td>− Keratosis obturans</td>
<td>− Malignant disease</td>
</tr>
<tr>
<td>− Foreign bodies – impacted</td>
<td>− Otitis media with effusion (rarely)</td>
</tr>
<tr>
<td>− Trauma</td>
<td>− Mastoiditis</td>
</tr>
<tr>
<td>− Malignant disease</td>
<td>− Otitis media with effusion (rarely)</td>
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<tr>
<td>• Dental</td>
<td>− Mastoiditis</td>
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<tr>
<td>– Caries</td>
<td>− Malignant disease</td>
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<td>– Abscess</td>
<td>− Otitis media with effusion (rarely)</td>
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<td>– Pericoronitis impacted</td>
<td>− Mastoiditis</td>
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<td>− wisdom toothl</td>
<td>− Mastoiditis</td>
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<td>– Costen's syndrome</td>
<td>− Mastoiditis</td>
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<tr>
<td>• Pharynx</td>
<td>− Mastoiditis</td>
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<tr>
<td>– Tonsillitis</td>
<td>− Mastoiditis</td>
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<tr>
<td>– Pharyngitis</td>
<td>− Mastoiditis</td>
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<tr>
<td>– Postoperative pain</td>
<td>− Mastoiditis</td>
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<tr>
<td>– Peritonsillar abscess</td>
<td>− Mastoiditis</td>
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<tr>
<td>− quinsy</td>
<td>− Mastoiditis</td>
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<tr>
<td>• Cervical spine</td>
<td>− Mastoiditis</td>
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<tr>
<td>– Osteoarthritis</td>
<td>− Mastoiditis</td>
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<tr>
<td>– Spondylosis</td>
<td>− Mastoiditis</td>
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<tr>
<td>• Abdominal</td>
<td>− Mastoiditis</td>
</tr>
<tr>
<td>– Mesenteric adenitis</td>
<td>− Mastoiditis</td>
</tr>
<tr>
<td>• Neurological</td>
<td>− Mastoiditis</td>
</tr>
<tr>
<td>– Herpes zoster (Ramsay Hunt syndrome)</td>
<td>− Mastoiditis</td>
</tr>
<tr>
<td>– Glossopharyngeal neuralgia</td>
<td>− Mastoiditis</td>
</tr>
<tr>
<td>− Trigeminal neuralgia (occasional)</td>
<td>− Mastoiditis</td>
</tr>
</tbody>
</table>

**Figure E.1** The nodular ulcerated appearance on the lateral margin of the pinna is typical of chondrodermatitis.

**Figure E.2** An extensive carcinoma of the pinna that had been neglected and brought the patient to treatment when he could no longer bear the pain.
EARDRUM

The external auditory meatus is highly sensitive, and even mild inflammation in the confined space of the meatus can lead to considerable pain. 

Foreign bodies may cause pain, usually after ineffective attempts at removal. Local infection of the skin and small furuncles may be extremely painful, the amount of pain bearing little relationship to the size of the furuncle, which may be only the size of a pin’s head. Impacted ear wax (or cerumen) is probably the most common ear problem in the general population. It may cause considerable pain and discomfort, especially following swimming and ineffective attempts at removal by syringing. Water may cause swelling of the wax, and sudden deafness follows complete occlusion of the meatus.

Further attempts at cleaning the ears lead to trauma, and a secondary otitis externa may develop. Diffuse otitis externa can arise as an acute episode or run a more chronic course. The auricle, and specifically the tragus, is tender on movement, and this is a valuable sign to differentiate the condition from other causes. In many cases, there is great swelling of the meatal skin, and it may not be possible to see the drum. The cause of otitis externa is often multifactorial. General conditions such as eczema, seborrhoeic dermatitis and neurodermatitis may predispose to infection with bacteria (the most common being Pseudomonas, Staphylococcus and Escherichia coli), fungus (Candida or Aspergillus) or viruses.

Malignant otitis externa, although not a neoplastic condition, may be a lethal condition in diabetic, aged and immunocompromised patients. It is caused by spreading bacterial or fungal infection of bone and subcranial tissues, thereby producing an osteitis or osteomyelitis of the skull base. This condition is common in hot and humid climates where it tends to be diagnosed early. Sadly, in more temperate climates, it is often overlooked, inadequately treated and consequently presents late and at an advanced stage with excruciating pain and facial palsy. Long-term antibiotic therapy, usually with ciprofloxacin at the outset, is required together with meticulous attention to correction of the predisposing condition.

MIDDLE EAR

Acute otitis media is one of the most common infections in childhood – hardly a child escapes one or more attacks. The infection usually accompanies upper respiratory tract infection, and the pain may develop within a matter of hours. It ceases abruptly when the drum ruptures and drainage is affected. The perforation usually closes spontaneously. Young children do not localize pain accurately, and earache for which there is no obvious local cause should prompt a wider search. Particularly aggressive acute otitis media may develop into acute mastoiditis. In this condition, the patient has an extremely high temperature (usually at least 39°C), and there is severe pain and tenderness behind and above the ear canal. Swelling and erythema develop in the postauricular sulcus, the pinna becomes proptosed, and the superior meatal skin sags. Early diagnosis and treatment is imperative as intracranial complications develop in those which are missed. Severe infection of the petrous apex may produce pain in the distribution of the ophthalmic division of the trigeminal nerve, as well as a lateral rectus palsy caused by inflammation of the abducens nerve (Gradengi's syndrome).

Chronic otitis media is usually painless but becomes painful during acute exacerbations of infection. The onset of pain in patients with pre-existing attico-antral or tubo-tympanic disease should cause concern as it often heralds spreading infection and imminent complications. Such complications include facial palsy, meningitis, brain abscess and sinus thrombosis.

Poor Eustachian tube function can be unmasked by the development of pain following rapid changes in atmospheric pressure. Most commonly, this is seen after diving or a prolonged air flight in a poorly pressurized cabin.

Secretory otitis media, so common in children, is usually characterized by a symptomless deafness. Dull aching pain or, rarely, stabs of acute pain may be a symptom. Malignant neoplasia of the middle ear and external canal is extremely uncommon, but when present is extremely painful. Other middle ear tumours (e.g. glomus tumours) are not associated with pain.

Referred pain

Referred pain affecting the ear is very common. Pain is referred through the Vth, VIIth, IXth and Xth cranial nerves, the upper cervical roots and the autonomic nerve supply. Despite careful examination, no abnormality is found in the ear.

Dental causes

Caries, dental abscess and impacted wisdom teeth are all common causes of earache. Malocclusion may give rise to dysfunction of the temporomandibular joint (Costen's syndrome). These patients often have a history of dental extractions or a badly fitting denture, and tenderness will be found over the joint especially on opening the jaw, movement that may well be limited.
Pharynx
*Tonsillitis* and *pharyngitis* are both sometimes accompanied by earache. Severe earache is almost always experienced about 1 week after tonsillectomy and should be controlled by non-steroidal anti-inflammatory analgesics. Frequent causes of referred pain from the pharynx are peritonsillar abscess, ulceration of the mucosa caused by trauma, and aphthae. The most important conditions from a diagnostic point of view are *malignant lesions* of the oro- and laryngopharynx. Earache is found very consistently in these conditions and may be the first symptom. Any patient with otalgia who has been a heavy smoker and for which there is no obvious local cause should have their pharynx examined very carefully.

Cervical spine
Cervical spondylosis affecting the C2 and C3 nerve roots may lead to referred pain around the ear.

Neurological causes
*Herpes zoster infection* of the geniculate ganglion (Ramsay Hunt syndrome) gives rise to intense otalgia associated with vesicular lesions of the external auditory canal and pinna. Facial palsy develops quickly, and may be permanent in up to 50 per cent of patients. The vesicular rash may also be seen on the palate and fauces. Rarely, hearing loss accompanies this condition.

In *glossopharyngeal neuralgia*, extremely severe, sharp pain is precipitated by swallowing. Pain that radiates from the throat to the ear may be due to glossopharyngeal neuralgia. Less severe variants can be associated with an elongated styloid process, which irritates the glossopharyngeal nerve when swallowing (Eagle’s syndrome).

**EATING, DISORDERS OF**

Andrew Hodgkiss
Disorders of eating are common and numerous. Both reduced and increased oral intake have a very wide differential diagnosis. This section is confined to a discussion of eating disorders as the term is generally used by psychiatrists and physicians, i.e. to a discussion of anorexia nervosa and bulimia nervosa, and the differential diagnosis relevant to each (Box E.2).

**Anorexia nervosa** is defined as a body weight maintained at least 15 per cent below that expected (or a body mass index of 17.5 or less), amenorrhoea in females, self-induced weight loss, distorted body image and a dread of fatness. Low body weight is usually achieved through a combination of reduced calorie intake and excessive exercise, although other strategies may be used. It is ten times more common in females than males, and the age of onset is typically in the mid teens. It is increasingly appreciated that those people reaching these criteria represent the tip of an iceberg of subclinical anorexia. Twenty per cent of 9-year-old girls in the UK have already tried dieting. Such subclinical presentations are sometimes dubbed ‘the anorexic stance’. Anorexia nervosa is overrepresented in the higher social classes and in risk occupations such as modelling, gymnastics and the performing arts.

Anorexia nervosa most commonly presents as unexplained weight loss in a young female. Other presentations include expressions of concern from relatives, primary or secondary amenorrhoea and abdominal pain. On examination, the severe anorexic has marked wasting, lanugo hair, a slow pulse, low blood pressure and cold extremities. Pubic and axillary hair is retained, differentiating anorexia nervosa from hypopituitarism, in which the patient also tends to be less emaciated. Other readily excluded causes of severe wasting are diabetes mellitus and thyrotoxicosis, but more difficulty may occur in distinguishing a malignancy, occult tuberculosis and malabsorption syndromes.

**Bulimia nervosa** is characterized by frequent episodes of eating large quantities of high-calorie food in a manner that feels subjectively out of control. Such binges are usually conducted in secret. The binge is usually followed by efforts to purge the body of the ‘forbidden’ food and so control the body weight despite the binge. Techniques include self-induced vomiting, laxative misuse, diuretics,
weight, either by not eating or by elimination through eating provokes anxiety that is relieved only by losing weight or losing weight. Any weight gain or that persists even when grossly fearful of becoming fat. There is a cardinal that always form the psychological backdrop in both but with the constellation of fears, urges and attitudes loss, dieting or any particular form of eating abuse, enter the differential diagnosis rests not with weight and all the physical and psychiatric conditions that enter the differential diagnosis. The Other psychiatric disorders, particularly depressive anorexia, enter the differential diagnosis. The distinction can sometimes be difficult, with marked weight loss, amenorrhoea, loss of energy, interests and concentration, sleep disturbance, obsessional ruminations and even suicidal ideation being potentially shared features. However, there are differences. Appetite is typically diminished in depression but normal or increased in eating disorders; more importantly, a preoccupation with food and its calorific content, a disturbance of body image and a phobia of normal weight are not features of a primary depressive illness. Similarly, these features distinguish an eating disorder from schizophrenia and obsessive-compulsive disorder, which may occasionally present with superficial resemblances through bizarre eating habits or food/weight-related rituals and overvalued ideas.

Indeed, the distinction between an eating disorder and all the physical and psychiatric conditions that enter the differential diagnosis rests not with weight loss, dieting or any particular form of eating abuse, but with the constellation of fears, urges and attitudes that always form the psychological backdrop in both male and female patients. There is a cardinal morbid fear of becoming fat that persists even when grossly underweight or losing weight. Any weight gain or eating provokes anxiety that is relieved only by losing weight, either by not eating or by elimination through vomiting, purgation or exercise. Food becomes an enemy, calories an obsession; life becomes dominated by the need to control this aspect of living. However, there may be great difficulty encountered in penetrating the patient’s resistance to elicit the information that is necessary to confirm the diagnosis. Failure to establish this psychological driving force after careful, repeated assessment should alert the physician to the possibility of one of the rare, cryptic presentations in the differential diagnosis.

**ENCOPRESIS**

Harold Ellis

(See also FAECES, INCONTINENCE OF, p. 193.)

Encopresis is often defined as the passage of faeces in socially unacceptable places; however, it is more accurate to restrict this term to the passage of normal stools in this way. Encopresis is therefore one of the forms of faecal incontinence in children, others being overflow soiling in association with a faecally loaded rectum (in children with non-Hirschsprung’s megarectum) or the incontinence associated with diarrhoea (especially if the stools are persistently loose, as in toddler’s diarrhoea).

Encopresis is normal in the first 3 years of life. Persistence of encopresis occurs with development delay, chaotic training or neglect, or neurogenic rectum (spina bifida, sacral agenesis, spinal tumours or idiopathically). Secondary encopresis (occurring after bowel control has been acquired) is most frequently a response to emotional stress, especially associated with anxiety or suppressed anger. Child physical or sexual abuse may present in this way.

The development of a neurogenic rectum due to spinal cords, cord damage from trauma or infection will lead to secondary encopresis.

**ENOPHTHALMOS (RETRACTION OF THE EYEBALL)**

Paul Carroll

Enophthalmos is defined as abnormal retrodisplacement (posterior displacement) of the eyeball. It may be a subtle clinical sign in which the palpebral aperture is narrower, and the upper eyelid may droop. This may occur in fractures of the orbital floor, where necrosis of the incarcerated orbital fat results in progressive enophthalmos. The enophthalmos that occurs in wasting diseases is due to the absorption of orbital fat, and the diagnosis as regards the eye presents no difficulty.
Paralysis of the cervical sympathetic nerves – Horner’s syndrome – can give rise to a mild degree of enophthalmos. Enophthalmos is always associated with the other well-defined symptoms of this lesion, namely, a diminution in the size of the palpebral aperture, constriction of the pupil, absence of sweating and blushing on the paralyzed side. Occasionally, it may be noticed that the hair over the affected side of the head is behaving differently from that on the unaffected side – it may lie flatter, or may lack lustre to a degree that the patient observes. The pupil is constricted, owing to the paralysis of the dilator fibres. In certain congenital cases (primary enophthalmos), there is well-marked retraction, associated with defective or irregular movements of the affected eyeball. Rarely, the condition is simulated by a maldevelopment of the globe, which has remained small, and is usually extremely hypermetropic and poor-sighted.

CAUSES OF ENOPHTHALMOS

- Congenital, for example microophthalmos (also known as nanophthalmos), due to abnormal development, and resulting in a small eye with abnormal function
- Phthisis bulbi, a description given to an atrophied eyeball with blindness and decreased intraocular pressure, which has occurred as a result of end-stage ocular disease, for example chronic endophthalmitis
- Fracture of the orbital floor: the soft tissue of the orbit prolapses, causing the eye to sink back into the orbit
- Atrophy of the orbital contents, secondary to irradiation of the orbit for malignant tumours, scleroderma or trauma
- Wasting diseases: HIV infection or malignancy
- Horner’s syndrome; cicatrising orbital lesions, such as metastatic sclerosing carcinoma and any chronic sclerosing inflammatory disease of the orbit

ENURESIS

Dipak Kanabar

(See also URINE, INCONTINENCE OF, p. 723.)

Enuresis means incontinence of urine in children, and it is usually nocturnal but can be diurnal. Children vary in the age at which they become reliably continent of urine. The majority of children are continent during the day by age 3 years, and at night by age 5. The prevalence of wetting diminishes with age, and approximately 15 per cent of 5-year-olds, and 3 per cent of 10-year-olds will wet the bed once a week or more. The problem is more common in boys than in girls (with a ratio of 2:1). There is a genetically determined delay in acquiring sphincter continence, with most children having a first-degree relative who was similarly affected.

Causes of incontinence include:

- An inherited trait: children whose parents used to wet the bed are more likely to do so themselves.
- A small bladder size.
- Infection: a urinary infection or cystitis.
- Constipation: faecal retention can also reduce bladder volume and cause bladder neck dysfunction.
- Low levels of antidiuretic hormone: children who wet the bed may have a lower level of this hormone, which suppresses the rate of urine production.
- Polyuria: this often leads to enuresis in children, so diseases such as diabetes mellitus, diabetes insipidus and chronic renal failure must be considered.
- Delayed growth and development: some children’s nervous systems are not mature enough to be able to sense when the bladder is full.
- Diet: dairy products, citrus fruits, chocolate and foods containing high levels of artificial colours and sweeteners have been connected with bed-wetting.
- Psychological and social factors: emotional problems can be a cause of bed-wetting. Secondary enuresis is common following family break-up or bereavement, and it is important to note that child sexual abuse may also present in this way.

The most common organic disease to lead to enuresis is urinary tract infection, especially in girls: enuretic girls have a fourfold increased likelihood of having a urinary tract infection compared with those without enuresis. Secondary enuresis (having previously been reliably continent) should be considered suspicious of a urinary tract infection, especially when associated with other symptoms such as urinary frequency, abdominal pain, dysuria and wetting in the daytime. Secondary enuresis can also be caused by neurological problems such as neurogenic bladder (associated with spina bifida or other cord lesions, such as tumours). Dribbling urinary incontinence may be the presenting sign of an ectopic ureter draining below the sphincter, or overflow from an obstructed bladder.

EPIDURAL

Reginald Daniel

Epiphora is the overflow of tears onto the cheek, and may be due either to increased production of tears or inadequate drainage of tears through the nasolacrimal drainage system (Box E.3).
INCREASED PRODUCTION OF TEARS

Psychic lacrimation is normally associated with pain or emotional upset. Reflex stimulation of lacrimation causing epiphora is commonly associated with irritative conditions or corneal disease. Corneal injury, a blast of air or a foreign body on the surface of the eye causes irritation of the trigeminal nerve that excites lacrimation. Even strong light, yawning, vomiting and laughing are associated with reflex lacrimation.

INADEQUATE DRAINAGE OF TEARS

This may be due to malposition of the lacrimal puncta, which should normally be closely applied to the eye in order to attract tears by capillary action. Such malposition may occur in ectropion associated with laxity of the lids, which may occur with age, or paralysis of the eyelid muscles that are responsible for blinking in facial palsy, or cicatricial lid disease, which pulls the punctum out of its correct position. Obstruction of the lacrimal drainage apparatus (canaliculus and nasolacrimal duct) will also cause epiphora, and this is in fact the most common cause of this problem.

EPISTAXIS

Michael Gleeson

It is important to realize that epistaxis (bleeding from the nose) is a sign, not a disease. Possible causes of the condition are shown in Box E.4.

Epistaxis often arises from the anterior part of the nasal septum (Little’s area). It is usually of a minor nature, and self-limiting. Bleeding from other parts of the nose may be extremely serious and, on occasion, life-threatening.

The nose receives its blood supply from branches of both the internal and external carotid arteries. The upper part of the nose is supplied by the anterior and posterior ethmoid arteries, branches of the ophthalmic artery (derived from the internal carotid), which enter it through the medial wall of the orbit. The rest of the nose receives its blood supply through the sphenopalatine branch of the maxillary artery and septal branches of the facial artery.

Very often, epistaxis can be controlled by simple measures, for example compression of the nostrils and application of cold packs. In childhood, when bleeding is almost always from Little’s area, these measures are usually effective. If this fails and the site of haemorrhage can be seen, bleeding can often be arrested by cautery. Otherwise, the nose has to be packed with nasal tampons that expand when irrigated with water. When packing with tampons is ineffective, 1 cm ribbon gauze impregnated with an antiseptic such as bismuth-iodoform paraffin paste may be used instead. All nasal packs should be carefully secured lest they fall backwards into the airway.

Continued bleeding despite local measures may require surgical or radiological intervention. Nowadays, endonasal endoscopic ligation of the sphenopalatine artery is possible and often effective. The sphenopalatine artery is found just underneath the posterior end of the middle turbinate in a bony canal where it can be clipped, diathermised or ligated. Other more radical approaches include intra-arterial embolization, or ligation of the external carotid artery in the neck, maxillary artery behind the maxillary antrum, and ethmoid arteries in the orbit.

Once the bleeding has been stopped and the patient’s condition is stable, attention must be devoted to identification of the cause. This is particularly indicated in patients with severe or
recurrent minor bleeds. A number of patients will be taking non-steroidal anti-inflammatory drugs, and these need to be stopped. Others may be anticoagulated, and this may need temporary adjustment. Blood film indices and coagulation studies should be carefully examined, to eliminate blood dyscrasias and coagulopathies. Hypertension alone rarely causes nose bleeds, but it is often an accompanying and exacerbating feature and should be controlled in the recovery phase.

It should be remembered that recurrent unilateral nose bleeds may be the first sign of a nasal tumour; if seen in an adolescent boy, they should alert one to the possibility of a juvenile angiofibroma (Figs E.3–E.5).

**ERECTILE DYSFUNCTION (IMPOTENCE)**

Paul Carroll

Erectile dysfunction (ED) or impotence are the terms used to describe the inability to achieve or sustain an erection for sexual activity to take place. It affects more than 1 in 10 men and has a number of causes, both physical and psychological. ED may be classified into mild, moderate and complete failure, based on a number of criteria including the level of sexual activity, frequency of full erections, and frequency of early morning erections on awakening. Ejaculatory insufficiency describes absent or reduced seminal emission. This may be because of a lack of external ejaculation or premature ejaculation and may be accompanied by lack of orgasm (Table E.1).

There is a progressive decline in sexual activity in association with a mild degree of hypogonadism in men with ageing. This decline is accelerated in the presence of ill-health. If hypogonadism is severe, usually associated with pathology, there is loss of libido and erectile function and a decrease in ejaculate volume. The majority of secondary cases of ED are caused by arterial occlusive disease. In subjects with diabetes mellitus, vascular abnormalities dominate in patients with type 2 diabetes; however, neurogenic factors play an important role in addition to vascular abnormalities in type 1 diabetes. The onset of ED in patients with hypertension is very frequently associated with the start of antihypertensive therapy. ED is a side effect of many of the commonly used

**Figure E.3** Magnetic resonance scan of a juvenile angioma (arrow). The patient, an adolescent male, presented with progressive, left-sided, nasal obstruction and torrential epistaxes.

**Figure E.4** At operation, the anterior end of the tumour is visible in the left nasal aperture.

**Figure E.5** The operative specimen. The tumour has been completely removed, albeit in two pieces.
antihypertensive agents including thiazide diuretics, beta-blockers, angiotensin-converting enzyme inhibitors and calcium-channel blockers. This side effect covers many classes of antihypertensive agent, which have varying mechanisms of action for lowering blood pressure. Therefore, the end result of lowering systemic blood pressure, which may interfere with maximal penile-filling capacity, rather than any individual effect unique to the class of antihypertensive, may be the causative factor in ED. Uraemia is associated with raised luteinizing hormone, oestrogen and prolactin levels, and low levels of testosterone. Approximately 50 per cent of males with uraemia suffer from ED. There is a high prevalence of ED among chronic alcoholics, who invariably have liver damage. The associated hypogonadism, feminization and neuropathy exacerbate the problem. Heavy cigarette smoking, in addition to its deleterious effects on peripheral vascular disease, is also associated with ED by the action of nicotine, which reduces corporeal blood flow and inhibits cavernosal venoconstriction. Peyronie’s disease, which starts off as an inflammatory process with cellular infiltrates followed by collagen deposition, is a cause of ED. The scars formed in this condition may distort the shape of the erect penis. The tunica albuginea is stiffened due to a depletion of elastic fibres and disables veno-occlusion. The plaque formation found at the periphery of the penis and in the midline interferes with arterial inflow. Both abnormalities of blood flow impair the tumescent and full erection phase for normal erectile function.

The prevalence of psychogenic ED is unclear. In many studies, psychogenic ED has been attributed to less than a third of cases. The measurement of this condition is compounded by the fact that complete vascular, neurological and psychological assessments are difficult to make in the non-erotic setting. In addition, abnormalities of nocturnal penile tumescence as a marker of ED are not a valid means of distinguishing among causes.

**EJACULATORY INSUFFICIENCY**

This condition is usually organic and is relatively common in neurological disease such as diabetic autonomic neuropathy, multiple sclerosis, paraplegia, etc. In the absence of any overt disorder, the two main causes of failure of ejaculation are retrograde ejaculation and anejaculation. The former can be diagnosed by the finding of many sperm in the post-coital urine sample. The latter may be difficult to diagnose, but there may be an absence of one or both vas deferentia and an absent or very small prostate.

A common problem is premature ejaculation. This may occur in younger men where the response to sexual stimuli may be very rapid. With reassurance from the doctor and familiarity with sexual stimuli, the problem usually resolves.

## ERYTHEMA

Barry Monk

Erythema – redness of the skin – is, along with swelling, heat and pain, one of the four cardinal signs of inflammation, and is the result of vasodilatation. There are many causes, both local and general (Box E.5).

**ERYTHRODERMA**

The term erythroderma is used to describe generalized redness of the skin; it is not a diagnosis in itself, as it may arise in psoriasis, eczema, drug reactions, or less commonly in cutaneous T-cell lymphoma. Erythrodermic patients cannot adequately control
heat regulation and may readily become hypothermic, or may collapse because of excessive loss of fluid through the skin.

LOCAL ERYTHEMA

A localized area of tender erythema may be the initial manifestation of an infection, such as a boil, or of cellulitis. The diagnosis usually becomes evident over the following hours, as the skin becomes rapidly hotter and more tender. Local infections in a limb may be associated with an ascending lymphangitis, manifested with a line streak of erythema moving towards the regional lymph nodes. Occasionally, malignant infiltration of the cutaneous lymphatics may present with linear erythematous streaks mimicking an infective lymphangitis, so-called lymphangitis carcinomatosa. In herpetic infections, the initial presentation may be with a local patch of erythema, the hallmark vesicles only becoming evident some hours later. Local erythema over a joint may suggest the diagnosis of gout. Erythema associated with swelling of the skin and intense itching occurs in an acute eczema, whether irritant or allergic in cause. Fixed drug eruption is an unusual disorder in which exposure to a causative drug (e.g. phenolphthalein, codeine or sulphonamides) gives a violent local erythema that rapidly darkens to a magenta colour, and usually blisters. It recurs at the same skin sites at each drug exposure.

SELECTED LOCALIZED PATTERNS OF ERYTHEMA

Rosacea

The characteristic feature of rosacea is a symmetrical erythema over the cheeks and nose, spreading onto the forehead and chin (Fig. E8). The affected area is red and swollen, and it is surmounted by sterile pustules and papules. The papules and pustules resolve with oral oxytetracycline, but the erythema tends to persist.

Palmar erythema

As an isolated phenomenon, this is best known in pregnancy, but it is also seen in chronic liver disease, thyrotoxicosis, rheumatoid arthritis and high-output cardiac states. It may also occur as a familial trait.

Erythema ab igne

This is a reticulate erythema that progresses to persistent haemosiderin pigmentation due to long-continued exposure to heat (infrared radiation). It used to be seen on the legs of elderly ladies with long-standing neglected hypothyroidism who sat close to open fires, but it is now most commonly encountered
on the abdomen in patients with chronic pancreatitis and other painful disorders who have used local heat as a form of analgesia.

**Erythema chronicum migrans (Lyme disease)**

The slowly spreading annular erythematous margin arising at the site of a tick bite is an important manifestation of Lyme disease (Fig. E.9). This disorder is now known to be a zoonosis, caused by the spirochaete *Borrelia burgdorferi*. The reservoir of infection is in deer, and the vector is the tick *Ixodes ricinus*. The condition is so named because of an outbreak of the disorder around the town of Lyme in the north-east of the USA. A rising specific IgA titre aids clinical diagnosis. Adequate treatment with penicillin or tetracycline is very important, as a proportion of untreated patients progress to serious central nervous system, cardiac or rheumatological problems, sometimes after the erythema has resolved.

Some very rare migratory erythemas can be cutaneous markers of underlying neoplasia; examples include necrolytic migratory erythema (glucagon-secreting pancreatic tumour) and erythema gyratum repens (carcinoma or lymphoma). 

**Erythema induratum (Bazin’s disease)**

This is a rare manifestation of tuberculosis that is seen in middle-aged women. The site affected is the posterior aspect of the legs, and the condition begins as a symmetrical eruption of deep-seated, painless red nodules. The surface becomes purple, and deep cold ulcers form, which have undermined borders. The patients show extreme sensitivity to tuberculin on skin testing.

**Erythema nodosum**

In this distinctive disorder, there is an acute eruption of hot, tender, erythematous nodules, usually over the extensor aspects of the lower legs, but the thighs, buttocks and extensor surfaces of the arms may
also be involved. The nodules are hard and deep, with shiny red overlying skin, but they gradually soften and their colour changes to violet and finally yellow; however, they never suppurate or ulcerate. An accompanying arthralgia is common, and there may be fever and malaise, whatever the underlying cause. Some cases are apparently idiopathic, but an assiduous search must be made for an underlying cause. The relative frequency of different causes will vary in different parts of the world. In Britain, the most common cause (among women) is sarcoidosis, but in patients of South Asian origin, tuberculosis should be suspected. Penicillin, sulphonamides and barbiturates are the most commonly implicated drugs. Rarer causes include ulcerative colitis, Crohn’s disease, histoplasmosis, coccidioidomycosis, blastomycosis, chlamydial infections and Behçet’s syndrome. A leprosy conversion reaction may manifest itself as an erythema nodosum-like phenomenon.

Erythema multiforme
This disorder is characterized by a sudden eruption of erythematous lesions arising especially over the hands, feet, buttocks and genitalia, often accompanied by lesions on the oral or genital mucosa and the conjunctiva (Fig. E.11). The skin lesions are classically polycyclic, giving the appearance of an archery target (‘target lesions’). In severe cases, the centre of the lesion may blister or become necrotic. When the mucosal lesions are severe, the condition is termed ‘Stevens–Johnson syndrome’. In some cases, no specific cause is found, but it may be triggered by an infection (herpes simplex, or Mycoplasma), a drug or radiotherapy. Spontaneous resolution occurs, but recurrent episodes may arise.

Toxic erythema
This is a widespread, symmetrical, blotchy erythema that tends to affect the trunk more than the extremities, and is often accompanied by malaise, fever and lymphadenopathy. This cutaneous reaction pattern may be provoked by many causes such as viral infections (especially glandular fever) and also drug hypersensitivity, especially to ampicillin, amoxycillin (Fig. E.12), sulphonamides and non-steroidal anti-inflammatory drugs. Spontaneous resolution, which usually occurs within 10–15 days, is often followed by desquamation.

Exanthems
The term exanthema is used to describe the erythematous rash that may arise in certain viral infections of childhood.

In measles, the rash usually develops on the fourth day, behind the ears. It spreads to the face and downwards to the trunk and extremities on the fifth to seventh days. The eruption is usually preceded by 1–7 days of prodromal coryza, nasal discharge and conjunctival injection, on the second or third day of which Koplik’s spots appear, as white specks surrounded by redness, on the buccal mucosa opposite the molar teeth. Koplik’s spots may be lentil-sized and few in number, or salt grain-sized and very numerous. The fever, which may have been high during the prodromal stage, persists while the rash appears and usually decreases as it fades. Cough, facial puffiness and photophobia are also diagnostic pointers.

Rubella usually affects an older child or young adult, and it is typically accompanied by posterior cervical and postauricular lymph node enlargement. The rash, which also begins on the face and spreads downwards to the trunk and extremities, is composed of faint pink
macules and fades in 3 days. Constitutional symptoms are mild, but joint pain may be prominent in adults. **Scarlet fever** usually affects children under 10 years of age, who become acutely ill with high fever and vomiting. The throat is red and oedematous, and there is characteristically a ‘strawberry tongue’. The rash, which is lobster-red with punctate deeper red lesions (likened to small spots of red ink on red blotting paper), appears on the second and third days behind the ears, spreading rapidly to the face, upper chest and flexor surfaces of the limbs. Purpura may develop in the skin creases. The fever persists with the rash from 2 to 5 days, and it is followed by desquamation in large flakes from the palms and soles.

**Erythema infectiosum** (fifth disease) affects children between 2 and 10 years of age. It begins with a ‘slapped-face’ erythema on the cheeks in an otherwise well child. Over the next few days, a maculopapular eruption spreads, sometimes in a gyrate pattern over the trunk and limbs. The illness is due to a parvovirus and heals in 1–2 weeks, although successive bouts may occur.

A macular erythematous rash accompanies the **seroconversion illness of primary infection with HIV** in around 50 per cent of patients. Between 10 and 14 days after exposure, fever, systemic toxicity and lymphadenopathy may occur. From 5 to 8 days later, oval macular erythematous lesions may appear on the trunk and limbs, extending beyond the usual T-shirt distribution of pityriasis rosea onto the palms and soles. The oral mucosa can be affected with superficial erosions. The whole illness most closely resembles secondary syphilis or glandular fever with rash. Lymphopenia and thrombocytopenia may occur. From 5 to 8 days after exposure, fever, systemic toxicity and lymphadenopathy may occur. From 5 to 8 days later, oval macular erythematous lesions may appear on the trunk and limbs, extending beyond the usual T-shirt distribution of pityriasis rosea onto the palms and soles. The oral mucosa can be affected with superficial erosions. The whole illness most closely resembles secondary syphilis or glandular fever with rash. Lymphopenia and thrombocytopenia may occur.

**Lupus erythematosus**

This is an important cause of erythema. It is more common in women than men and is usually classified into two varieties: systemic and discoid. The systemic form affects joints, kidneys and the haemopoietic, cardiovascular, respiratory and central nervous systems, as well as the skin, and it occurs in a younger age group than the discoid form, which is a dermatological disorder. In the skin, both types of lupus erythematosus cause perifollicular inflammation that is followed by scarring. The classic lesion of lupus erythematosus is an atrophic, red, scaly plaque with follicular plugging. The histopathology is characteristic, showing epidermal thinning and basal cell liquefaction. Direct immunofluorescence microscopy demonstrates linear staining at the basement membrane in the lesions of both types.

**Systemic lupus erythematosus (SLE)** should be suspected in young women presenting with weakness, fever, weight loss, arthralgia and proteinuria, as well as a persistent erythematous rash on the face or areas exposed to sunlight. The so-called ‘buttery rash’ over the face, although well known, occurs in only a minority of patients with cutaneous manifestations of SLE. Although typical lupus erythematosus plaques may be found – particularly over the dorsa of the hands and fingers – they are less common than other erythematous rashes seen in SLE, such as urticaria, urticated plaques, livedo reticularis, photosensitivity and nailfold erythema. A diffuse telogen effluvium is not infrequent. Established cases often have tell-tale signs on the hands; as well as nailfold erythema and telangectasia, there may be infarcts of the cuticles, finger nodules and pulp atrophy. Some 80–90 per cent of such patients will have circulating antinuclear antibodies in high titre and raised DNA-binding proteins. Different varieties of antinuclear antibody exist and relate to differing clinical presentations, and this is the subject of current clinical research. The antinuclear antibodies can often be demonstrated throughout the skin (e.g. by taking a biopsy of non-light-exposed normal skin). This test is negative in the discoid variety. A systemic pattern, but without renal mortality, can be induced by certain drugs, such as hydralazine, procainamide, griseofulvin and phenytoin.

**Chronic discoid lupus erythematosus** lesions largely occur on sun-exposed sites such as the cheeks, forehead, nose and perioral skin, although the scalp is not infrequently affected. Lesions persist for many months, extending slowly, and leaving depigmented, central atrophic and hairless scars. Close examination of the erythematous active edge shows telangectasia and follicular plugging (which can be demonstrated by detaching an adherent scale and observing downward-projecting ‘tin-tack’ plugs). Although the direct immunofluorescence test of such lesions will be positive, in 80 per cent of patients no circulating antinuclear antibodies are found. Less than 5 per cent of patients progress to the systemic form.

**Dermatomyositis**

The principal cutaneous signs of dermatomyositis are erythema and oedema (Fig. E.13). The proximal myopathy is described in detail elsewhere. There is often marked discordance between the severity of skin and muscle disease in any individual patient. The dermatological hallmark is a particular periorbital bluish-red oedema (‘heliotrope’ erythema): periungual changes identical to those of SLE (see above) may be seen, often with papules over the dorsa of the fingers.
A diffuse scaling erythema of sun-exposed areas can occur, as well as psoriasiform lesions over the elbows and knees. In common with SLE, there is histological basal cell liquefaction and follicular plugging. A patient over 40 years of age with dermatomyositis should be investigated for underlying neoplasia, which will be present in about 40 per cent of cases. Childhood dermatomyositis is almost never associated with malignancy, but it can be responsible for considerable and disabling soft tissue calcification.

EUPHORIA

Andrew Hodgkiss

Euphoria describes an elevation of a person’s mood. It is frequently the presenting symptom of bipolar affective disorder (manic depression) or schizophrenia, but these diagnoses should not be made before drug-induced mania and organic mood syndromes have been ruled out. The causes of euphoria are listed in Box E.6. Mania is one of the most common and most clearly identified causes of euphoria, but the diagnosis is frequently missed or delayed until the patient and family have come to harm through a combination of grandiosity and poor judgement. Diagnosis is based on the presence of either elated or irritable mood associated with some or all of the following features: overactivity; pressure of speech; racing thoughts or flights of ideas; distractibility; grandiose ideas that may be delusional; decreased sleep; decreased appetite; and behaviour that indicates poor judgement. The patient may become sexually disinhibited, spend too much money, or drive recklessly, endangering life. The mood can have the quality of infectious good humour, but irritability is also common, and at interview euphoria may be quickly and dramatically replaced by tearfulness, feelings of depression and remorse lasting for a few minutes before euphoria once again dominates the picture (labile mood).

In bipolar disorder, bouts of euphoria lasting more than a week or so may be preceded or followed by periods of depression. Cyclothymia describes a milder chronic form of bipolar illness in which the subject is prone to constant swings between mood elevation and depression, with only brief stretches of normality. In such cases, a moderate level of euphoria persisting for weeks or even months serves the subject well by increasing drive and enthusiasm, leading to increased sociability and a greater involvement in sexual, political, religious or occupational activities. When a patient’s mood is significantly elevated, it can be a difficult judgement for the clinician to decide when to try to intervene. Mania sometimes presents for the first time in the puerperium. In some depressed patients, it is precipitated by the use of treatments including antidepressants or electroconvulsant therapy. The acute phase of a severe manic illness may be impossible to distinguish from schizophrenia, and the diagnosis will only be clarified by follow-up. In schizophrenia itself, states of excitement with increased activity and euphoria are quite common, although often there is a bizarre quality to the thoughts and incongruity of affect that distinguishes the schizophrenic states from manic illness. Schizoaffective disorder describes those
conditions which have features both of schizophrenia and depression or mania.

Many organic conditions may present with an elevated mood resembling a manic episode, and these include causes of delirium (see p. 133) in which there will be impairment of consciousness. Hyperthyroidism may produce a manic syndrome, and the use of corticosteroids, L-dopa, bromocriptine and the anticholinergic drugs procyclidine and benzhexol may cause euphoria.

Damage to the frontal lobes is sometimes associated with mild euphoria out of keeping with the patient's situation. The mood is often facile or facetious and accompanied by changes in personality, including disinhibition, deteriorating social behaviour and a loss of spontaneity and volition. Slow-growing frontal tumours (e.g. meningiomas), multiple sclerosis and dementia (Alzheimer's and Pick's diseases) may cause a frontal lobe syndrome. Temporal lobe epilepsy may be associated with a psychosis resembling manic illness. Rarely, neurosyphilis and porphyria can cause euphoric or grandiose states.

Many drugs are taken because of their ability to cause euphoria, the most common being alcohol. The list of compounds used to obtain a 'high' include sympathomimetics (amphetamine, dextroamphetamine, methamphetamine, methylphenidate and appetite suppressants); cannabis; cocaine; hallucinogens such as LSD, mescaline and magic mushrooms; inhalants including the aliphatic and aromatic hydrocarbons found in petrol, glue, paint and typewriter correction fluid; spray-can propellants; anaesthetic gases, for example nitrous oxide and ether; short-acting vasodilators such as amyl or butyl nitrite; opioids (heroin and morphine) and many compounds prescribed as analgesics, anaesthetics or cough suppressants, such as codeine and methadone; and sedatives, hypnotics and anxiolytics, for example benzodiazepines and barbiturates. The diagnosis of a drug-induced euphoric state is made from the history and circumstances of the presentation, and by urine drug-screening for substances. Typically, the altered mood state is relatively brief. Diagnostic difficulties can arise in patients with a history of prolonged drug abuse, for example with cocaine, which may trigger a psychosis indistinguishable from mania or schizophrenia. The picture is further complicated because patients with bipolar disorder or schizophrenia may be heavy drinkers or drug misusers, and they may experience an exacerbation of their underlying condition through the drugs they are using.

**EXOPHTHALMOS (PROPTOSIS)**

Paul Carroll

Exophthalmos (also called exophthalmia) is anterior protrusion of the eye from the orbit. This may be bilateral or unilateral.

**BILATERAL EXOPHTHALMOS**

The most common cause of this condition is Graves’ disease, in which the exophthalmos is associated with thyroid gland swelling and other general symptoms of thyrotoxicosis. The degree of prominence of the eyes is variable, in some cases being so great that there is inadequate lid coverage of the cornea on attempted eye closure. A protrusion causes retraction of the upper eyelid; consequently, the eyes look wide open, giving the patient an expression of alarm or astonishment (Stellwag’s sign; Fig. E.14). When the eyes are lowered, the upper lids lag behind the downwards excursion of the eye, leaving a broad portion of the sclera visible above the cornea (von Graefe’s sign). The extent of exophthalmos may be asymmetrical, with minimal involvement of the contralateral eye. The condition sometimes appears to be unilateral. Increasing oedema of the lids along with inflammation of the conjunctiva and dilatation of the vessels over the insertion, particularly of the lateral rectus, are significant findings. The myopathy of Graves’ disease most frequently involves the vertically acting muscles, with limitation of upward eye movement.

Bilateral exophthalmos is also associated with:

- Chronic obstructive pulmonary disease
- Superior vena cava syndrome
- Cushing’s syndrome

Other uncommon causes of bilateral exophthalmos are septic thrombosis of the cavernous sinus, historically

![Figure E.14 Stellwag’s sign. Reproduced with kind permission of the patient.](image-url)
associated with skin infection at the inner angle of the eye or ethmoidal sinus suppuration; bilateral lymphomatous deposits; pseudotumour; and, in children, the craniodysostoses.

UNILATERAL EXOPHTHALMOS

Unilateral exophthalmos may be due to:

- Orbital cellulitis
- Orbital neoplasms
- Meningioma
- Lymphoma
- Lacrimal gland tumour
- Optic nerve glioma
- Pseudotumour (a space-filling lesion with the characteristics of a chronic infiltrating inflammatory process)
- Carotid–cavernous sinus fistulae
- Cavernous sinus thrombosis
- Cavernous haemangioma
- Mucocoele
- Infiltrative disorders (e.g. sarcoid, Wegener’s granulomatosis and xanthogranuloma)

Orbital cellulitis can begin as a primary inflammatory process in front of the orbital septum, thereafter extending backwards or rising from direct orbital extension of a paranasal sinus infection (Fig. E.15). The dire complication of further progression is septic cavernous sinus thrombosis. The general signs and symptoms of cavernous sinus thrombosis are more serious than in uncomplicated orbital cellulitis. Headache, nausea, vomiting and altered consciousness are early signs. Venous congestion produces gross chemosis and proptosis with a bluish discoloration of the eyelids. There is early onset of pan-ocular motor paresis. Bilateral involvement is virtually diagnostic of cavernous sinus thrombosis. Pseudotumour involvement of the orbit usually first appears above, but may also be found in, the inferior retrobulbar tissues. Another area of involvement is at the apex of the orbit, when visual loss may be of early onset. A pseudotumour along the superior orbital fissure causes a painful external ophthalmoplegia (Tolosa–Hunt syndrome).

Capillary haemangioma is the most common cause of unilateral proptosis in childhood. Adult cavernous haemangioma causes a slowly progressive exophthalmos.

Orbital lymphomas usually occur in the anterior orbit and involve the conjunctiva and eyelids. However, posterior orbital involvement may occur, and it is important to exclude a primary systemic lymphoma. Lacrimal gland tumours occur characteristically at the upper lateral portion of the orbit, causing painless downward and medial displacement of the globe and irregular enlargement of the lacrimal fossa.

Dermoid cysts specifically appear in childhood at the upper lateral quadrant of the orbit. Superior and medial swellings are more suggestive of mucocoeles of the frontal or ethmoidal sinus. Teratomas may occur congenitally or in very early life. Rhabdomyosarcoma is the most common intraorbital malignant tumour of childhood, presenting within the first decade of life. The development of proptosis is rapid, occurring within 1–3 weeks. The rapidity of progression is more or less pathognomonic of this tumour.

Optic nerve gliomas usually manifest before the age of 5 years and result in a downwards nasal and forwards proptosis. Meningiomas usually occur in older women, but they can occur in childhood, where their course is much more rapid. Orbital haemorrhage may arise from trauma or sudden extreme physical effort and cause bleeding from orbital varices, leading to alarming, progressive orbital swelling. Arteriovenous fistula can occur following a fracture of the base of the skull, with rupture of the internal carotid artery as it passes through the cavernous sinus. The carotid–cavernous sinus fistula causes a pulsatile proptosis of the eyeball associated with a bruit that is synchronous with the pulse. Gross dilatation of the conjunctival vessels is visible, with arterIALIZATION of the conjunctival veins. Intermittent unilateral exophthalmos in children following coughing or crying is nearly always associated with a deep cavernous haemangioma.

Ultrasound scanning, computed tomography, magnetic resonance imaging and biopsy are all useful in assessing cases of exophthalmos.
EYE, BLINDNESS OF
Reginald Daniel

(See also VISION, DEFECTS OF, p. 739.)

The World Health Organization defines blindness as a central visual acuity of less than 3/60 (1/20), or a visual field of less than 10 degrees. An alternative functional definition is loss of vision sufficient to prevent one from being self-supporting in an occupation, making the individual dependent on other persons, agencies and devices in order to live.

Colour blindness is a genetically determined disorder that is a minor handicap and is not true blindness.

The causes and prevalence of blindness through the world vary from country to country. It is estimated that 75 per cent of blindness in the world is avoidable.

The leading causes of blindness are trachoma, leprosy, onchocerciasis, xerophthalmia and cataract. In Western countries, age-related macular degeneration, diabetic retinopathy and glaucoma are the most common problems.

Different categories of those who are blind have different needs, and the agencies for the blind assess the individual blind person’s requirements and provide a variety of services, including mobility training, visual magnifying aids, talking books, training in Braille, educational assessment, job rehabilitation and psychological counselling.

EYE, INFLAMMATION OF (RED EYE)
Reginald Daniel

Inflammation of the eye may involve the conjunctiva (as a conjunctivitis), the cornea (keratitis, usually in the form of a corneal ulcer) and, less commonly, the uvea (uveitis) and sclera (scleritis). Localized patches of episcleritis may superficially resemble conjunctivitis, and the dusky circumcorneal congestion in an acute glaucoma may simulate that of an acute anterior uveitis. The character of the inflammation varies with the type of the disease, but certain symptoms such as pain, photophobia and lacrimation are common to all inflammatory conditions, and are by themselves of little value in the differential diagnosis.

CONJUNCTIVITIS
In conjunctivitis, the conjunctival vessels are dilated; they are freely movable over the subjacent sclera, and the conjunctival injection is most evident at a little distance from the corneal margin. The circumcorneal portion of the conjunctiva, owing to its firmer attachment to the sclera in this region, is relatively less injected. If the condition is purely conjunctival, the cornea is clear and bright, the anterior chamber and iris are normal in appearance, and the pupil is black with normal reactions. Purulent discharge may occur, and there is often a feeling of grittiness as of sand or dust in the eye.

The main types of conjunctivitis are infectious (bacterial, viral or fungal), allergic, giant papillary and reactive.

Bacterial conjunctivitis (Fig. E.16) commonly results from infection with Staphylococcus aureus, Streptococcus pneumoniae, Haemophilus influenzae, Pseudomonas aeruginosa and Chlamydia trachomatis. This type of conjunctivitis generally commences in one eye and can spread to both, often causing a heavy, thick, mucopurulent ocular discharge. The lids may be stuck together after sleep, and the pain is generally slight and allayed by closing the eyes.

In trachoma, which is a chlamydial infection endemic in the Middle East but rare in the UK, the conjunctiva is studded with enlarged follicles, particularly on the undersurface of the upper lid and in the upper conjunctival fornix. The follicular enlargement is associated with thickening and oedema of the tissues of the upper lid and in the upper conjunctival fornix. The follicular enlargement is associated with thickening and oedema of the tissues of the upper lid, causing partial ptosis, with excess lacrimation and, in the later stages, vascular infiltration (pannus) of the upper part of the cornea. In the later stages of trachoma, the infiltration is followed by scarring, which may distort the tarsal plate, leading to cicatral entropion and trichiasis (Fig. E.17).

In ophthalmia neonatorum (acute conjunctivitis of the newborn), caused by infection from the birth canal (Chlamydia, Staphylococcus or gonococcus), there is often a profuse mucopurulent discharge. The condition is differentiated from imperfect canalization of the nasolacrimal ducts by the fact that, in the latter,
the discharge is present without the accompanying inflammation. Untreated cases are at grave risk of secondary corneal ulceration and require identification and early antibiotic therapy.

Viral conjunctivitis usually produces a watery discharge. Typically, the infection starts in one eye and quickly spreads to the other. Viral conjunctivitis is self-limiting and usually lasts several weeks. The viruses that commonly cause conjunctivitis include the adenovirus, herpes simplex, Coxsackie, rubella and rubeola viruses, and even the common cold or influenza virus. The eyes may feel uncomfortable, and there may be tender lymphadenopathy around the ear or neck. Typical cold-like symptoms of viral infection may be experienced, such as a sore throat and runny nose. As the infection is self-limiting, medication is not usually prescribed, but for the worst cases topical steroid therapy may be prescribed to reduce the discomfort from the inflammation.

Fungal conjunctivitis can result from fungi such as Candida albicans, occurring more commonly in immunocompromised individuals, such as HIV-positive patients or those receiving immunosuppressant therapy.

Allergy-based conjunctivitis usually affects both eyes and causes much irritation and tearing. The eyelids are often swollen, as is the conjunctiva (chemosis), and frequently a whitish, stringy discharge occurs. There is usually an urge to rub the eyes. The airborne allergens include pollen, the house-dust mite and mould spores. Allergic conjunctivitis occurs commonly in individuals with other allergies, such as hayfever, asthma and eczema. The chemicals used in eyedrops, particularly the preservatives, frequently result in allergic conjunctivitis. There is a particular type of allergic conjunctivitis termed ‘vernal conjunctivitis’, which is a bilateral recurrent condition that tends to occur in children during warm weather and goes into remission during the cooler months. The palpebral conjunctiva develops large flat-topped conjunctival nodules (giant papillae) resembling cobblestones. Giant papillary conjunctivitis is a conjunctivitis most commonly associated with contact lens use. Patients complain of increased mucus formation and mild ocular irritation on removing their contact lenses. They suffer with increased awareness of the lenses, blurring of vision after prolonged lens wear and eventual contact lens intolerance. The mechanism by which giant papillary conjunctivitis occurs is not clearly understood, but it seems to be related to the contact lens materials and preparations used to care for the lenses (cleaning and sterilization solutions). An immune system response and overwear of the lenses, especially worn or soiled lenses, are other probable factors.

Reactive conjunctivitis occurs due to a chemical, an irritant or a foreign body in the eye. Noxious fumes, air pollution, chlorine in swimming pools, acids or alkalis can result in this form of conjunctivitis. Trauma, including surgical, can result in reactive conjunctivitis with resulting conjunctival hyperaemia and tearing, as can entropion (inturning of the lower eyelid), which allows the eyelashes to rub against the cornea and conjunctiva (trichiasis). Strong ultraviolet irradiation damages the conjunctiva and corneal epithelium when using a sun-lamp or arc-welder without eye protection. This can cause reactive conjunctivitis, producing conjunctival hyperaemia with considerable irritation or pain.

KERATITIS

In keratitis, there is loss of clarity of the corneal tissue, with the hyperaemia most marked near the corneoscleral margin. Keratitis is inflammation of the cornea, which can result from a variety of bacterial, viral or fungal infections, or may be non-infective, for example due to trauma or associated with an autoimmune disease. Inflammation of the cornea commonly presents as a painful red eye with reduced visual acuity due to cellular infiltration and later corneal oedema. Discharge is usually present and may be watery, mucoid or purulent. The pupil may be small due to reflexive miosis, and photophobia is common. Fluorescein dye readily demonstrates any ulceration (an epithelial breach). Interruption of an intact corneal epithelium permits the entrance of microorganisms into the corneal stroma, where they may proliferate and cause ulceration. Corneal ulcers produce greyish or white opacities of the cornea.
stroma with loss of the corneal epithelium. In more serious untreated cases, infiltrations of the cornea may lead to a loss of corneal tissue and progress to perforation of the cornea. In severe cases, there may be pus in the anterior chamber – termed a 'hypopyon' (see Fig. C.26, p. 112). The diagnosis presents no difficulty; the ulcers are obvious if the cornea is examined carefully and stained with fluorescein.

IRITIS
In iritis or 'anterior uveitis', the eye is congested and painful (in contrast to a 'posterior uveitis', or choroiditis, which simply blurs the vision). This vasodilatation differs from that in a conjunctivitis in that it is most evident in the circumcorneal region, with the tarsal conjunctiva remaining unaffected, and that the colour of the injection is brick-red rather than pink. The cornea retains its clarity, but the aqueous humour may be turbid due to the presence of cells and protein, and there may be punctate deposits of leucocytes on the posterior surface of the cornea (keratic precipitates), or rarely a hypopyon or pus level within the anterior chamber (see Fig. C.25). Owing to the increased vascularity of the iris, and to the exudation into iris substance, its volume is increased and its mobility impaired; hence the pupil becomes small and sluggish. The presence of blood and exudate in the substance of the iris also changes its apparent colour – a blue iris becomes greenish – and the fine detail of the iris structure is blurred and obliterated. Adhesions are apt to occur between the iris and the lens at the point of their immediate contact, the edge of the pupil; in the constricted state of the pupil, these may not be seen. On dilatation with cyclopentolate or atropine, these adhesions or posterior synechiae prevent the enlargement of the pupil at certain points, and it therefore becomes irregular in shape (Fig. E.18).

GLAUCOMA
Acute-angle closure glaucoma is a disease of the later years of life, and of hypermetropes rather than myopes. It is precipitated by any of the factors that may provoke dilatation of the pupil. At first, the chief complaint in subacute attacks is of temporary obscuring of vision and the appearance of haloes or rainbows around light sources. There is also often a feeling of tension in the eye and a dull frontal headache in addition to the loss of vision. In acute attacks, the pain is severe, radiates from the eye to the head, the ears and the teeth, and is associated with nausea – a symptom that may lead to the mistaken diagnosis of migraine. The lids may be oedematous and the conjunctiva injected (Fig. E.19). The cornea is hazy due to oedema, the anterior chamber shallow, the iris appears discoloured, and the pupil is mid-dilated and fixed. The eye is hard to the touch and very tender. Vision fails rapidly, even down to bare perception of light, within a few hours. The distinction between subacute or acute glaucoma (as just described) and chronic simple glaucoma

![Figure E.18 Iridocyclitis synechiae. (Moorfields Eye Hospital.)](image1)

![Figure E.19 Acute glaucoma. (Moorfields Eye Hospital.)](image2)
is easily made. Chronic simple glaucoma is an asymptomatic disease, and is usually discovered in the course of routine examination. No pain, blurring of vision, haloes or feeling of tension are complained of, and the visual field loss that characterizes this disease is rarely noticed by the patient in the early stages.

The importance of discriminating between iritis and acute-angle closure glaucoma cannot be overemphasized; the use of atropine or some similar mydriatic is a basic treatment of iritis, while in acute glaucoma it is disastrous (Table E.2).

Acute inflammation of the eye may be seen in episcleritis and scleritis (Fig. E.20). Episcleritis may be either simple or nodular, producing localized injection of the episcleral vessels. In either type, the condition is normally idiopathic and asymptomatic, with the patient merely complaining of redness of the eye. Scleritis is a more serious condition, often associated with the connective tissue disorders. In contrast to episcleritis, the eye is painful and tender, with a deep-seated bluish injection. Recurrent episodes of inflammation of the sclera may produce progressive scleral thinning. Early treatment with systemic anti-inflammatory drugs or corticosteroids is mandatory.

Table E.2 A summary of the points of distinction between conjunctivitis, iritis, acute glaucoma and keratitis

<table>
<thead>
<tr>
<th></th>
<th>Conjunctivitis</th>
<th>Iritis</th>
<th>Acute glaucoma</th>
<th>Keratitis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Conjunctiva</td>
<td>Conjunctival vessels bright red and injected; movable over subjacent sclera; injection most marked away from corneoscleral margin; colour fades on pressure</td>
<td>Ciliary vessels injected, deep red; most marked at corneoscleral margin; colour does not fade on pressure</td>
<td>Both conjunctival and ciliary vessels injected but dusky in colour</td>
<td>Conjunctival vessels red and injected; injection most marked near the corneoscleral margin</td>
</tr>
<tr>
<td>Cornea</td>
<td>Clear, sensitive</td>
<td>Clear, sensitive</td>
<td>Steamy, hazy, insensitive</td>
<td>Irregular reflex, corneal opacity</td>
</tr>
<tr>
<td>Anterior chamber</td>
<td>Clear; normal depth</td>
<td>Aqueous may be turbid</td>
<td>Very shallow</td>
<td>Normal or possibly hypopyon</td>
</tr>
<tr>
<td>Iris</td>
<td>Normal</td>
<td>Swollen, may be adherent to lens and muddy coloured</td>
<td>Injected</td>
<td>Normal</td>
</tr>
<tr>
<td>Pupil</td>
<td>Black, active (normal)</td>
<td>Small and fixed, later festooned after adhesions form to lens</td>
<td>Mid-dilated, fixed, oval</td>
<td>Black active (normal)</td>
</tr>
<tr>
<td>Intraocular tension</td>
<td>Normal</td>
<td>Normal</td>
<td>Raised</td>
<td>Normal tension</td>
</tr>
</tbody>
</table>

Figure E.20 Episcleritis. [Moorfields Eye Hospital.]
The patient’s features, expression and facial movements can in many cases suggest an instant diagnosis. Although such ‘spot’ diagnoses may often be wildly inaccurate, in many cases they prove more telling than many investigative procedures performed later to prove or disprove a diagnosis. Experience alone can teach the student to detect all that is to be learned from the patient’s facies. The more subtle abnormalities of expression, the play of the emotions, and the response of the features to questioning and intellectual and emotional challenge are transient and fleeting, and cannot be recorded or reproduced. Indeed, they are sometimes so intangible as to defy any attempt to describe them. The passive vacant aspect of someone who is a chronic alcoholic and the tremor of their mouth when they open it to protest their temperance are clinical observations that cannot be reproduced visually, except by a television camera. The shifty eyes of the drug addict, the fatuous placidity of the patient with advanced multiple sclerosis, the anxious look of those within a few days of death, the explosive suddenness with which the victim of multiple sclerosis or brain damage bursts into laughter or tears, the vacant stare of the child with learning difficulties, the unsmiling sad appearance of someone who is depressed, the distant removed look of the schizoid personality, the excessive vivaciousness of the hypomanic personality – these are a few of the many familiar and striking lessons of the face that must be seen in real life if they are to be learned and utilized. It is upon the appearance of the face that people most rely for the judgement of general health and well-being, for this is the only part of the body that everybody is habitually accustomed to see – plumpness or wasting, the complexion, the expression, the carriage of the head, the way the eyes, brows, cheeks and mouth move, for example, may all suggest certain disorders. Appearances may be deceptive, however. Pallor is by no means the same thing as anaemia; a ruddy complexion is not necessarily a sign of rude health; it is often far from easy to distinguish the appearance of illness from the expression of unhappiness; it is all too easy to mistake for aggression what is really shyness.

**MYXEDEMATOUS FACIES**

The skin of the myxoedematous face is coarse, dry, sallow, pale and waxy, with occasionally a tinted rose-purple flush over each cheek (Fig. F.1). The puffiness of the eyelids may suggest acute glomerulonephritis, but the subcutaneous tissue everywhere is of firm consistency, and doughy rather than oedematous. The tongue is enlarged. The nose is broadened, the ears are thickened, and the lips are swollen. The hair is scanty, receding from the forehead, the eyebrows are thin and sparse (although the scantiness of the outer half often regarded as a diagnostic feature occurs too frequently in normal subjects for this to be reliable), and the nails are brittle and striated. Masses of fatty tissue may be found in the neck and trunk. The slow, husky speech, the expressionless face and the general attitude of the patient may superficially suggest parkinsonism, but the diagnosis may be made by paying attention to other clinical features.

In hypopituitarism, the eyelids and nose – in contrast to myxoedema – are unaffected and show no undue thickening. Another point of differentiation is that, in pituitary disease, complete loss of axillary and pubic hair is common – a feature that does not always occur in myxoedema. In hypopituitarism, the face is hairless in males or females and unduly wrinkled. The features are those of a middle-aged Peter (or Pauline) Pan. In contrast, the patient with myxoedema looks like a wax doll who has been left in a sunlit shop window for too long. In hypopituitarism, the skin is soft and smooth and the hair of soft texture, whereas in myxoedema the hairs and skin are of coarser quality. The voice in myxoedema is a husky croak, but in hypopituitarism it is normal.

![Figure F.1](image-url) Facies in myxoedema (a) before and (b) after treatment. [Dr P. M. F. Bishop.]
CONGENITAL SYPHILITIC FACIES
The victims of congenital syphilis, now an extreme rarity, may present a facies that is unmistakable – an overhanging forehead, perhaps frontal bosses, a depressed nasal bridge, striated scars radiating from the corners and other parts of the lips, with a sallow, earthy complexion. Closer observation of the eyes and teeth may detect the opacities of old keratitis, as well as the changes in the upper incisors that were stated by Jonathan Hutchinson to be pathognomonic. These teeth are wide-gapped, irregular, and so deficient in enamel over the anterior and median parts of their cutting edges that the resulting crescentic notch imparts a striking appearance.

MYOPATHIC FACIES
Many cases of myopathy show no characteristic facies; in facioscapulohumeral dystrophy, the face is always involved, the muscles around the mouth being affected most with a loose pout of the lips at rest and ‘transverse’ smile (tire en travers). These features are due to defective facial musculature, particularly to weakness of the orbicularis oris. Paresis of the orbiculares palpebrarum is often evident when an attempt is made to close the eyes, although it may sometimes lead to prominent and perhaps staring eyeballs. An inability of the patient to whistle or to blow out their cheeks demonstrates the weakness of the orbicularis oris, which is often rendered obvious by the large amount of labial mucous membrane exposed while the mouth is at rest.

In dystrophia myotonica (myotonia atrophica, Steinert’s disease), ptosis, facial weakness and dysarthria occur. There is a characteristic weakness and diminution in the size of the sternomastoids, and the masticatory muscles are poorly developed or waste early, giving a long, lean facial appearance. In males, frontal baldness is common. In the rare ocular myopathy, there is progressive ptosis of the eyelids and immobility of the eyes.

MYASTHENIC FACIES
In patients suffering from myasthenia gravis, there are two types of facies. The first is the patient whose lids lag with fatigue. The second depends on the characteristic myasthenic smile, almost a sneer. This unfortunate and misleading facial expression is the result of deficient action on the part of the zygomatic and risorius muscles, and it exemplifies the curious way in which in this disease some muscles are affected and others escape, even when they derive their innervation from the same source. In patients with this disorder, facial weakness is worsened by repeated movement, but it responds rapidly (albeit transiently) to anticholinesterase drugs such as intravenous edrophonium, which acts more rapidly than neostigmine. In ocular myasthenia, only the extraocular muscles are involved.

HYPERTHYROIDISM
The facies of hyperthyroidism depends chiefly upon the ‘stare’ (see Fig. T.13). Surprise or terror is suggested by the prominence of the eyeballs and the retraction of the eyelids. The degree of exophthalmos varies greatly, and it may be completely absent; it is sometimes unilateral. The sclera is visible between the edge of the iris and the eyelids; the usual harmony of movement between the eyeball and the eyelid is lacking; and normal blinking is much diminished or entirely in abeyance. The surface of the conjunctiva may be abnormally bright and glistening, and the secretion of tears may be excessive. In contrast with the white of the eyeballs, there is often considerable dark pigmentation of the eyelids, which may also be the site of some oedema. The size of the pupils varies, with undue dilatation occurring only in exceptional cases. The upper eyelid lags as the eye follows the examiner’s finger downwards. Eye movements are often diminished in range due to intrinsic muscle weakness, and the muscles of the brow are wasted, giving diminished wrinkling on raising the eyebrows (Joffroy’s sign). A moist skin and a readiness to flush may often be noted in the face.

PARKINSONISM
In this disease, a cardinal symptom is muscular rigidity that affects the skeletal muscles generally, as well as those of the face. However, the ocular muscles escape and, as a consequence, while the face as a whole is expressionless or ‘mask-like’, the eyes appear to move with natural or even abnormal rapidity. For instance, they will turn in the direction to which the patient desires to look before the head has assumed a corresponding position. The face often has a staring expression, the eyelids being retracted by the tonic spasm of the orbiculares palpebrarum. An absence of normal blinking has been ascribed to the same cause. In contrast with the slow development of facial expression, there may under the influence of emotion be marked want of control over the fully developed emotional movement, and the patient protests that the exuberance of their laughter or tears is entirely out of proportion to their feelings of merriment or sorrow. The poverty of facial and general movement may falsely suggest a lack of intelligence and mental activity.
FACETIC FACIES

In a considerable number of the few remaining cases of tabes dorsalis, the appearance of the face is sufficiently striking to afford a clue to diagnosis. The small size or the inequality of the pupils reacting to accommodation but not to light (Argyll Robertson pupils) may first attract attention. The drooping of the upper eyelids, combined with some wrinkling of the forehead produced by a compensating effort on the part of the frontalis muscle, imparts a sad expression. This drooping of the eyelid is not due to any paresis of the levator palpebrae superioris – as may be shown by the raising of the lid when the patient is looking upwards. Rather, it depends on the fact that this muscle – like most muscles of the body – is in a condition of hypotonia, so that under the influence of gravity the lid hangs like a half-raised curtain in front of the eyeball. In other respects, the face may be normal, but the majority of tabetics have a sallow complexion and very little subcutaneous fat – two conditions that contribute to their generally unhealthy appearance. Many victims of this disease exhibit a deficiency of the emotional reflex movements of the facial muscles; during conversation, the play of the features appropriate to the subject of their talking is not so noticeable as in the case of healthy individuals.

FACIES OF ACROMEGALY

In acromegaly, changes in appearance frequently take place to such a degree that the patient becomes unrecognizable to friends who have known him or her only before the onset of the disease. These are the result of abnormal growth of the bony and subcutaneous tissues, especially in the skull and extremities. The characteristic facies is brought about by osseous hyperplasia of the frontal ridges, the mastoid, zygomatic, malar and nasal processes, while the lower jaw is usually enlarged in all directions. The prominent, arched brows, with retreating and wrinkled forehead, the massive nose, the long, thick upper lip and the heavy chin (Fig. F.2) form the most conspicuous features. The lower teeth are unduly wide apart, and may project some distance in front of the upper. The tongue may be so enlarged as to keep the mouth open and to display many fissures and indentations as the result of its pressure against the teeth. In some cases, the lower jaw is not affected, and the face may be described as abnormally square.

DOWN’S SYNDROME

This facies is so distinctive that the diagnosis may usually be made at a glance. The head is brachycephalic; the palpebral fissures slant obliquely inwards and downwards towards a broad flat nose, rendered even broader by the presence of epicanthus; the eyelids show signs of chronic blepharitis; the ears are large and pitcher-shaped; the lips are fissured and often left open to allow a coarse tongue to protrude; the forehead is downy, and the hair of the scalp scanty, wiry and frequently mouse-coloured; and the complexion is florid and mottled. The almond-shaped eyes, the presence of epicanthus, the florid complexion

Figure F.2 A case of acromegaly, exemplifying the heavy enlargement of the front lower jaw. Before and after treatment. Reproduced with kind permission of the patient.
and the absence of fatty masses serve to distinguish Down’s syndrome from cretinism.

HEPATOLENTICULAR DEGENERATION (WILSON’S DISEASE)
The characteristic facies of this disease is seen only in advanced cases and may be described as one of ‘fixed emotion’. The slightest attempt to engage in conversation may evoke a sustained expression of exaggerated mirth, which is quite unlike that seen in other diseases of the nervous system. There is also a tendency to fall to one side when in the sitting position. The illness is associated with bilateral degenerative changes in the lenticular nuclei, together with cirrhosis of the liver, due to the excessive amounts of copper in the tissues. The most remarkable feature of the disease is the Kayser–Fleischer ring, which is present in about 50 per cent of patients; this is a ring of rusty-brown pigment at the periphery of the cornea. Radiating brownish spokes of copper carbonate on the anterior or posterior lens capsule less often cause the characteristic ‘sunflower’ cataract.

MITRAL STENOSIS
It is occasionally possible to suspect mitral stenosis at sight, on account of the remarkable malar hyperaemia and dark-crimson lips contrasting with the yellowish pallor of the forehead, perioral and perinasal skin. If one covers the malar regions and the lips, the face looks sallow, yet the malar flush and the dark-crimson lips give a look almost of plethora. When cardiac failure occurs and the liver becomes engorged, an element of icterus may be added.

PRIMARY POLYCYTHAEMIA
The coloration of the nose, lips, ears and palpebral conjunctiva is the chief feature of the facies in this malady, presenting an appearance that may be described as a combination of exposure to weather, of plethora and of cyanosis. The diagnosis depends on discovering pronounced polycythaemia, and generally a large, firm spleen. Polycythaemia may also be secondary to other conditions, cardiac, pulmonary or malignant disease.

CIRRHOSIS OF THE LIVER
There is nothing characteristic in the facies when cirrhosis of the liver is at an early stage. Nor can one diagnose the existence of cirrhosis with certainty even when the facies is that of chronic alcoholism, with its telangiectases over the cheeks, coarsening of the tissues (especially on and around the nose and mouth), and purplish reddening in general. However, in the later stages of cirrhosis, the sallow, dull, diffusely pigmented facies is often distinctive, although the actual peculiarities are not easily described.

ACUTE GLOMERULONEPHRITIS
The generally swollen, half-bloated look and the partial closing-up of the eyes by oedema are usually unmistakable, but a somewhat similar appearance may be presented by the effects of insect bites, with angioneurotic oedema, or after the administration of aspirin or other drugs to which the patient is allergic. In the nephrotic syndrome (Fig. F3), kwashiorkor and many other conditions with extensive oedema, this may extend to involve the face, although it does not usually do so in cardiac or starvation oedema, which affects the dependent parts. In leprosy, a similar puffy appearance of the face may be seen, particularly around the eyes; a variety of skin lesions may occur, with nodules, plaques and thickening of the skin. The ear lobes may enlarge, and the lines of the face may coarsen and become deeper, giving the so-called ‘leonine facies’. Patches of depigmentation may occur, or there may be bronzed hyperpigmentation. The eyebrows and eyelashes may fall out, and the lips swell. Nasal blockage may occur and a saddle-nose deformity may develop.

ARTHRITIS AND CONNECTIVE TISSUE DISORDERS
In dermatomyositis, the most characteristic rash consists of a dusky red eruption on the face, over the nose and cheeks, periorbital regions, occasionally on the forehead and on the neck, shoulders, front and back of the chest and arms. The erythema may be mottled or diffuse, and either intensely red or cyanotic, or a mixture of both. Sometimes, a dusky lilac hue is seen on the upper eyelids – the so-called

Figure F.3 Facies resulting from nephrotic syndrome due to cancer.
‘heliotrope rash’ – which is said to be typical of dermatomyositis. Telangiectasia may be present, as it may in systemic lupus erythematosus and scleroderma.

A common skin lesion of *lupus erythematosus* has a ‘butterfly’ distribution over the bridge of the nose and the cheeks (Fig. F.4). The facial skin lesion of *sarcoidosis* may have the same distribution, but the eyelids and ears may be infiltrated with brownish nodules.

In *scleroderma*, the parchment-like skin may be so tightly drawn over the underlying muscles that the face becomes completely expressionless (Fig. F.5) and the mouth cannot open widely.

In *giant-cell (temporal) arteritis*, the inflamed temporal arteries are tender to touch and become thrombosed. Vision may be affected if the retinal arteries become involved. Tophi in the ears indicate the presence of *gout*.

**ACANTHOSIS NIGRICANS**

The outstanding feature of this disease is the extreme pigmentation that develops in various parts of the body, as in the axillae, groin, nipples and umbilicus, but also in the neck or face. The degree may be described as what would, more or less, result if a collier’s hands were stroked over the skin, producing massive darkening, almost blackening, in the areas affected. Although rarely generalized, it is usually bilateral and symmetrical, blending into adjacent normal skin. Although the disease usually indicates abdominal carcinoma (especially carcinoma of the stomach), the patient may present for treatment only on account of the pigmentation, without any suggestion at the time that there is malignant disease anywhere. It is probably an extreme degree of the liability to diffuse pigmentation of the skin that malignant disease in general tends to produce. It may precede malignant disease or follow it, but usually appears at the same time. It may also occur in Cushing’s syndrome, acromegaly, Stein–Leventhal syndrome, suprarenal insufficiency, pituitary or hypothalamic tumours, or other lesions at the base of the brain. When associated with a malignant process, these tend to be aggressive, and rapidly fatal. Neither clinically nor histologically can acanthosis nigricans associated with a malignant process be differentiated from the disease without this association.

Other colorations of diagnostic significance are those of *haemochromatosis*, in which over 90 per cent of patients show bronzing of the skin from melanin deposition, about half having haemosiderin deposition.
also causing a slate grey colour. The bluish tinge of the cartilage of the ears and sclerae in ochronosis appears usually between the age of 20 and 30 years. The cartilages of the ears may be slate-blue or grey, and are often thickened and irregular. Pigmentation of the sclera is usually localized to a small area halfway between the cornea and the inner or outer canthus. The skin over the malar areas and nose is often darker than usual. Other abnormal colours that may be seen in the face include the patchy pigmentation of the chloasma of pregnancy, or that of vitiligo or albinism, and that resulting from the prolonged administration of arsenic.

**ADDISON’S DISEASE**
Generalized darkening of the skin of the face may be the first thing to attract attention in a case of Addison’s disease, but the distinctive character of the pigmentation is that it occurs in the mucous membranes within the mouth (Fig. F.6), where it tends to be grey, as well as on the skin of the face and other parts of the body, where it is dark brown.

**CUSHING’S SYNDROME**
The red ‘moon face’ and hirsutism (see HIRSUTISM, p. 280) are characteristic, and are often seen as the result of corticosteroid therapy. Although the features look plethoric, there is no true polycythaemia. This is one point of differentiation from the features of simple obesity, others being the presence of bruising (ecchymoses), muscle weakness, wide purple–red striae (those in simple obesity being more narrow and pink) and hypertension. In Cushing’s syndrome, the cheeks may become so chubby that, when full face, they obscure the ears (Fig. F.7).

**ARGYRIA**
This condition is rare nowadays, but it may still be met with among those working with silver. The coloration is even and uniform; there is a blue–grey appearance that persists when pressure is applied to the skin, which does not blanch as does a cyanotic skin. It is a subcutaneous rather than a dermal pigmentation. The features in pachydermoperiostosis, a rare familial condition associated with pseudohypertrophic osteoarthropathy (with finger clubbing), are typical with thickening and furrowing of the face, deep nasolabial folds, greasy skin of the face and scalp, and often excessive sweating. The condition appears to be transmitted by an autosomal dominant gene with variable expression.

**ACUTE ILLNESS**
Erysipelas, measles, scarlatina and mumps often permit an immediate facial diagnosis. Cellulitis also is self-evident. In lobar pneumonia, the bright eyes, flushed cheeks, active alae nasi and labial herpes constitute what may fairly be termed a ‘typical picture’. Respiratory distress advertises itself by expression of anxiety and fear in pulmonary and cardiac disease, although alterations in colour due to cyanosis contribute to the appearance. Labial herpes (herpes febrilis) may also accompany many other febrile diseases, even a simple coryza, and may be due to sun sensitivity. Herpes zoster may affect the face and the peri-orbital region (Fig. F.8).

**ALTERATIONS IN CONTOUR**
Slight facial asymmetry is very common. Marked asymmetry occurs in patients with lipodystrophy, hemiatrophy or hemi-hypertrophy, or congenital absence of the condyle of the mandible. Lack of teeth or bad dentures may contribute to asymmetry, as may swelling of the parotid or other salivary glands.
or of the lymph nodes. Some rarer conditions may be mentioned as generally identifiable at sight. In osteitis deformans (Paget’s disease), the face has the shape of an inverted triangle and, in consequence of the prominence of the forehead, appears to be toppling forwards. In leontiasis ossea, there is progressive irregular enlargement of the bones of the cranium and face, with consequent asymmetry; the superior maxilla is particularly prominent. The rare condition of oxycephaly (‘steeple head’) need be seen only once to be subsequently recognizable.

THE EYES
The eyes alone often provide diagnostic evidence of general as well as local disease. Pigmentation, oedema of the lids and exophthalmos have been mentioned. A squint may demand a detailed consideration of the central nervous system, as will spontaneous nystagmus. Icterus of the conjunctivae may be evidence of hepatic disease, and the comparatively rare but striking appearance of blue sclerotics points to fragilitas ossium. ‘Bags under the eyes’ (‘steeple head’) need be seen only once to be subsequently recognizable.

VOLUNTARY MOVEMENTS
Weakness of the facial muscles is discussed in FACE, PARALYSIS OF (p. 184).

Abnormal movement of the jaw may be due to any painful condition of the temporomandibular joint.

INVOLUNTARY MOVEMENTS
Besides the tremor of the head of old age and parkinsonism, tremor may be due to alcohol, tobacco or other drugs. There is also a familial tremor of the hands, face and/or head affecting several members of the same family, usually commencing before the age of 25 years. The head-nodding of children may be mentioned in this connection. Other involuntary movements point to chorea, which may be hereditary (Huntington’s), rheumatic or (rarely) senile, to habit spasms or to tics (see p. 100). In aortic regurgitation, there may be a constant jerking of the head that is synchronous with the heart beat (De Musset’s sign). Facial paralysis and the peculiar condition of facial hemiatrophy or hemi-hypertrophy are sometimes obvious, but sometimes evident only on careful examination.

EXPRESSION
Patients’ expressions at interview may provide an indication of their attitude not only to their illness, but also to their physician and what is expected of him or her. Differentiation of the emotional from the physical factors may be very difficult. There may be an expression of melancholy or depression, of anxiety, nervous tension or querulousness. In some cases, depression hangs over the patient like a black cloud, the face being dull, without hope for the future, expressionless and uninterested in what is going on around.

FACE, PAIN IN
David Werring & Mark Kinirons

Pain in the face may be due to: (i) local disorders of the sinuses, soft tissues, joints and teeth; (ii) neurological or neurovascular causes; and (iii) atypical facial pain (Box F.1). Here, the emphasis will be on neurological and neurovascular causes, but the other causes will be briefly discussed.

LOCAL DISORDERS

Ears, nose, sinuses and teeth
Sinusitis causes pain and tenderness localized to the frontal or maxillary sinuses, and percussion over the affected area worsens the pain. Blowing the nose and bending forwards also worsens the discomfort. The nose will often be blocked. If simple measures, including decongestants and postural drainage, are ineffective, referral to an ear, nose and throat specialist is recommended. Nasopharyngeal carcinoma frequently infiltrates the cranial nerves, especially the trigeminal nerve, causing facial pain. Disease of the petrous
The pain usually starts in either the maxillary
pathology
syndromes)
SALIVARY GLAND DISEASE
NEUROLOGICAL/NEUROVASCULAR
TRIGEMINAL NEURALGIA
GLOSSOPHARYNGEAL NEURALGIA
CLUSTER HEADACHE
SUNCT (short-lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing)
PAROXYSMAL HEMICRANIA
MIGRAINE
TEMPORAL ARTERITIS
POST-HEPATIC NEURALGIA
RAMSAY HUNT SYNDROME
TOLOSA–HUNT SYNDROME
ATYPICAL FACIAL PAIN AND HEMIFACIAL SPASM

Box F.1 Causes of facial pain

• Ears, nose and throat disease
• Sinus disease
• Teeth disorders
• Local diseases (muscles, ligaments, soft tissues, sinuses)
  – Temporomandibular joint pathology
  – Muscle tension (myofascial syndromes)
  – Salivary gland disease
• Neurological/neurovascular
  – Trigeminal neuralgia
  – Glossopharyngeal neuralgia
  – Cluster headache
  – SUNCT (short-lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing)

Temporal bone infection or neoplasm causes pain in the first trigeminal division and ipsilateral abducens palsy (Gradenigo's syndrome). Dental caries or root canal sepsis can cause pain in the first and second divisions of the trigeminal nerve, which is worsened by hot or cold liquids. Malocclusion of the teeth has been reported to cause trigeminal neuralgia-type pain (see below).

Temporal mandibular joint disease
Dysfunction of the temporomandibular joint, also termed facial arthromyalgia, is poorly understood. It has been suggested to be due to malocclusion or loss of the molar teeth. The main clinical findings are tenderness of the temporomandibular joint and muscles of mastication, trismus (jaw spasm), limited or jerky jaw movements, and evidence of bruxism (tooth-grinding) or frictional damage to the buccal mucosa and the tongue. Signs of subluxation of the joint include clicking noises and lateral displacement of the meniscus. Pain in the temple, face and neck with a sensation of ear fullness constitute the rather doubtful nosological entity of Costen’s syndrome.

Neurological and neurovascular causes

Trigeminal neuralgia
Trigeminal neuralgia almost always begins after the age of 40, except when associated with multiple sclerosis. It is more common in women than men (ratio 3:2). Trigeminal neuralgia is characterized by paroxysmal facial pain that is intense and described as ‘shooting’, ‘jabbing’, ‘stabbing’ or ‘like an electric shock’. Each paroxysm lasts under 30 seconds, with less than a minute between each successive pain. Each series of paroxysms lasts a few seconds to a few minutes. The pain usually starts in either the maxillary or the mandibular division (or both), and rarely in the ophthalmic division, to which it may later spread, and is provoked by stimulation of a trigger point in the same division as the pain. Contact with the trigger point may occur during washing, shaving, combing the hair, blowing the nose, talking or eating, or even from a slight draught on the face. Patients typically describe techniques to avoid touching the face, by contrast with other causes of facial pain, in which massage is typically helpful. The pain is nearly always unilateral. The severity of the pain often causes the patient to screw up the affected side of the face in a grimace, hence the name ‘tic douloureux’. Before effective treatments were available, suicide was said to be common. The pain tends in the early stages of the illness to occur in bouts lasting days or weeks. The periods of pain subsequently become longer with shorter remissions, so that eventually, although still intermittent, the pain will occur on most days. On examination, there are no abnormal signs in the nervous system, except in association with multiple sclerosis, where sensory loss in the trigeminal distribution may be found. Trigeminal neuralgia occurs in about 3 per cent of patients with multiple sclerosis.

The pain may be relieved in more than half of the patients by the drug of first choice, carbamazepine. For patients who do not respond to or tolerate carbamazepine, oxcarbazepine is a good alternative. For those refractory to or intolerant of carbamazepine and oxcarbazepine, baclofen may be tried instead. Lamotrigine, phenytoin and gabapentin may also be effective.

For patients who still have significant and distressing symptoms despite medical therapy, it is reasonable to discuss surgical options using microvascular decompression, rhizotomy, or gamma knife radiosurgery. The type of surgical procedure offered varies widely in different hospitals, and there are few high-quality data on efficacy. Radio-frequency or glycerol ablation of the Gasserian ganglion can be performed, although the 5-year recurrence rate is over 50 per cent. An aberrant vascular loop is now, with the increasing use of magnetic resonance imaging, recognized as a cause of trigeminal neuralgia, and in these cases surgical microvascular decompression can be effective. Gamma-knife treatment to the trigeminal nerve has also recently been used. Relapse may occur following medical or surgical treatments, and the prognosis is difficult to predict with any certainty. Spontaneous remissions frequently occur early in the illness.
Glossopharyngeal neuralgia
This condition is much less common than trigeminal neuralgia. It causes pain that has a similar lancinating character, but which is localized instead to the ear, base of tongue or jaw angle. Pharyngeal and otalgic variants have been described, depending on where the pain occurs. Glossopharyngeal neuralgia is triggered by talking, swallowing or coughing. It may be idiopathic or secondary to compression of the nerve by a tumour, infection or aberrant blood vessels. Glossopharyngeal and trigeminal neuralgias may co-exist, and the medical treatments are similar.

Cluster headache, migraine and cranial arteritis
Cluster headache and the other trigeminal autonomic cephalgias, migraine and temporal arteritis are conventionally considered to be mainly headache disorders (see HEADACHE, p. 258).

Migraine
Migraine is fully considered in the section on HEADACHE, p. 258.

Post-herpetic neuralgia and Ramsay Hunt syndrome
Herpes zoster is caused by reactivation of the varicella-zoster virus, which lies dormant in the trigeminal, geniculate and dorsal root ganglia following chickenpox infection in infancy or childhood. The vesicular rash is commonly in the trigeminal distribution, most often the ophthalmic division. Less commonly, the external auditory meatus or upper cervical roots are affected. A herpetic rash in the external auditory meatus and a facial palsy constitute the Ramsay Hunt syndrome and are due to involvement of the geniculate ganglion. Pain in the ear or trigeminal distribution may precede the rash, or appear without any rash, causing diagnostic difficulty. Post-herpetic neuralgia is pain persisting beyond a month after the rash has crusted over. The pain may be burning and distressing, with misperception of light touch stimuli as painful over the affected area (allodynia). Symptomatic treatment with standard neuropathic pain drugs (e.g. carbamazepine and gabapentin) may be helpful.

Atypical facial pain
Atypical facial pain is not a clear diagnostic entity but refers to a poorly understood group of conditions that are quite common in both medical and surgical practice. Atypical facial pain is commonly considered a diagnosis of exclusion. Minor surgery or injury to the face, teeth or gums may initiate symptoms, but these persist without a clear local cause. Nevertheless, there do seem to be some characteristic features. The pain is often poorly localized to a deep, non-muscular, non-neuralgic distribution in the face. It is usually deep, boring, aching, dragging or nagging in quality, and it is long-lasting, from weeks to years, without relief. It quite often starts following one or more dental procedures, and patients may subsequently be referred to a number of specialties (ear, nose and throat, maxillofacial and neurology) in the course of their illness. The pain may be worsened by fatigue or stress. It may be associated with symptoms of anxiety or depression, and may respond to antidepressant treatment, particularly with tricyclic drugs. Examination is normal, and there are no helpful investigations.
Upper motor neurone facial paralysis occurs in any condition affecting the corticofacial projections in the cerebral hemisphere. In practice, stroke, cerebral mass lesions and inflammatory conditions (e.g. multiple sclerosis) are the most likely causes. Because the descending corticospinal motor fibres become closely packed together as they descend, in lesions involving the lower corona radiata or internal capsule, facial weakness is usually associated with arm or leg weakness. Facial weakness may be associated with aphasia in dominant hemisphere lesions. Isolated upper motor neurone facial weakness has been reported with very small lacunar infarctions in the internal capsule or corona radiata.

LOWER MOTOR NEURONE PARALYSIS (PERIPHERAL FACIAL PALSY)

This condition occurs as a result of a lesion of the VIIth nerve nucleus or of the nerve itself (Fig. F.9). The upper and lower halves of the face are affected equally, and there is no dissociation of emotional and voluntary movements. If there is no recovery, contractures may occur, the corners of the mouth being drawn to the affected side, thereby giving a false impression of weakness on the normal side. **Facial myokymia** is a fine rippling movement of all muscles on one side of the face. It is an important physical sign, as it is most often seen in cases of multiple sclerosis and brainstem glioma, and if noted makes brain imaging (ideally with magnetic resonance imaging) mandatory. The causes of lower motor neurone facial palsy are listed in Box F.2.

By far the most common form of peripheral facial palsy is **Bell’s palsy**, described by and named after the British surgeon Charles Bell in 1821. The onset is rapid, and the patient often awakens in the morning to find the face paralysed on one side; in other cases, the condition takes a day or two to develop. In about one-half of patients, there is a dull pain or numbness just below the mastoid and behind or in front of the ear at the onset. The majority will have a change in auditory acuity (hyperacusis, in which sounds seem louder) due to involvement of the nerve to stapedius. Some patients will have disturbed taste in the anterior two-thirds of the tongue, or impaired salivary flow due to involvement of the chorda tympani. There may be decreased lacrimation (tearing) ipsilateral to the lesion if it is proximal to the geniculate ganglion. The eye cannot be closed fully and is liable to injury by dust. On examination, there is weakness of the facial muscles and platysma, so that the face is pulled away from the affected side on smiling etc. On attempted eye closure, the globe deviates upward and inward (Bell’s phenomenon). A careful search should be made for vesicles in the external auditory meatus (which suggests herpes zoster of the geniculate ganglion; Ramsay Hunt syndrome) and for facial myokymia (which suggests pontine glioma or demyelination).

The cause of Bell’s palsy is not known, although 60 per cent of patients have a preceding viral infection, and there is increasing evidence that herpes simplex virus

**Figure F.9** Bilateral lower VIIth nerve palsy.
FACE, SWELLING OF

Harold Ellis

In this section are included only swellings of the skin and subcutaneous tissues. Malignant and other diseases of the facial bones, etc., are considered under JAW, SWELLING OF (p. 314) and SALIVARY GLANDS, SWELLING OF (p. 596). It is necessary therefore to determine the anatomical site of the lesion before considering the pathology. For example, swelling of the parotid gland will lie below and in front of the ear, or in the anterior prolongation of the gland, lying on the outer surface of the masseter. Swelling of the sublingual gland will be seen in the floor of the mouth close to the frenum, while lateral to this will be felt the submandibular salivary gland, which is also palpable from outside in the submandibular fossa. Suspected swellings in either the sublingual or submandibular gland or its duct should therefore be palpated bimanually, between a finger below the angle of the jaw and the finger in the floor of the mouth.

Occasionally, a patient may present with painless symmetrical oedema of the face, commonly of the eyelids, where the tissues are loosest. This will almost certainly be of renal origin, since cardiac oedema causes oedema primarily in the dependent parts. Another form of oedema that may involve the whole face, but chiefly the eyelids and lips, is angioneurotic oedema. The recurrent attacks, each of sudden onset, the familial history, the associated symptoms of burning and irritation and the presence of similar areas in other parts of the body should clinch the diagnosis.

Swelling of the face and neck is seen in Cushing's syndrome, whether primary or secondary to corticosteroid therapy, and when there is obstruction to the venous return to the heart from the head and neck, as is seen with mediastinal and bronchial neoplasms. In trichiniasis, oedema of the eyelids is common, although more diffuse oedema of the face may occur.

A well-defined cystic swelling on the face is most commonly a sebaceous cyst, a structure that is freely movable on the deeper tissues but attached to the skin. A dermoid cyst is much rarer, and occurs only at lines of suture, the most common site being above the outer canthus of the eye (external angular dermoid).

A cyst in this situation is strongly suggestive of a dermoid origin; the diagnosis is confirmed if there is attachment to bone but not to skin, and particularly if depression of the bone has occurred (as it does in long-standing cases), the edge of the depressed area being palpable. Meningocoele may occur occasionally as a translucent swelling at the root of the nose. It will be present at birth and will exhibit an impulse on coughing or straining. Haemangiomas are frequently found on the face, and may appear cystic on palpation; however, their dusky colour and surrounding dilated vessels will give the clue to their identity. They also empty on pressure. Pigmented naevi will be recognized on sight.

Type I infection of the nerve is the main pathogenic agent. An immune mechanism is possible, given the association of Bell's palsy with interferon-alpha-2b treatment and the lymphocytic infiltrate seen on histology. The prognosis for recovery is excellent: 80–90 per cent of patients recover fully by 4–6 months. The elderly, and those with poorly controlled hypertension, seem to have a less favourable prognosis. The main long-term complications, occurring at 3–4 months, are contractures and synkinesis of the face.

Treatment involves careful eye protection (lubricating drops, taping at night and consideration of an eye patch if there are signs of corneal damage). Corticosteroids are probably helpful, but must be given within 72 hours, before irreversible neuronal denervation has occurred. Treatment with approximately 1 mg/kg daily in divided doses (e.g. 25 mg twice a day) for 10 days is a suggested regimen. High-dose steroids should not be stopped abruptly as rebound worsening may occur. Current evidence does not demonstrate benefit from using antiviral agents, although in severe Bell's palsy they can be considered as an adjunct to steroids.

Other causes of lower motor neurone facial paralysis are less common than Bell's palsy, and are listed in Box F2. Most of these are associated with other symptoms and physical signs, which should provide clues to the correct diagnosis. For example, in Lyme disease, a history of visiting endemic areas (e.g. the New Forest in the UK) and a migrating erythematous rash may be found, whereas Guillain–Barré syndrome in its full form includes limb paralysis, glove and stocking sensory symptoms and areflexia. In cases of tumours at the cerebellopontine angle, ipsilateral stockineurosyndrome, whether primary or secondary to corticosteroid therapy, and when there is obstruction to the venous return to the heart from the head and neck, as is seen with mediastinal and bronchial neoplasms. In trichiniasis, oedema of the eyelids is common, although more diffuse oedema of the face may occur.

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Solid tumours of the face are lipomas and fibromas. The latter are fairly common and include an important variety, the neurofibromas. These tumours vary in size from being quite minute to 2–3 cm or more in diameter, and they may be hard or soft. Other stigmas of von Recklinghausen’s disease such as pigmentation (either diffuse or in multiple café-au-lait spots), or a profusion of soft, fleshy neurofibromas in other parts of the body (chiefly the trunk) help in the diagnosis. The condition sometimes runs in families.

Rodent ulcer is particularly common on the face and eyelids; it is the exception to find it elsewhere. It starts as a small nodule, often with a ‘pearly’ appearance, but soon breaks down to form the characteristic indurated ulcer with hard, rolled edges (Fig. F.10). Epithelioma, with its raised everted margin and indurated base, and possibly secondary enlargement of regional nodes, is another malignant condition found on the face, particularly the lips (Fig. F.11). Confusion may arise in distinguishing epithelioma from the innocent condition molluscum sebaceum. However, molluscum runs a short course and the centre sloughs, leaving an unsightly scar. Biopsy must be performed early in any suspicious lesion.

Various inflammatory swellings are found on the face, of which the following are some of the most important:

- **Boils and carbuncles** are common, particularly around the lips. They have the same character as elsewhere, except that oedema is more marked.
- **Erysipelas** is prone to occur on the face. It is marked by a vivid red oedematous swelling associated with fever. The redness tends to spread, the edges being raised and well defined from the healthy skin. The oedema may be continuous, or it may disappear in one place and reappear in another. In very severe cases, the fever is high, rigors occur, the cuticle may be raised in blebs, and sloughing may ensue.
- **Alveolar abscess** and **dental caries** are fertile sources of facial swelling, as is an abscess in the nasal sinuses (see JAW, SWELLING OF, p. 314).
- **Anthrax** chiefly affects operatives in wool and horsehair factories and workers of raw hides. The disease is characterized by the formation of a vesicle, which bursts, forms a scab, and then becomes surrounded by a ring of vesicles around which is an area of oedema. The diagnosis is confirmed by discovering anthrax bacilli in the discharge; a fluid prepared from a drop of fluid from one of the vesicles contains long chains of large, square-ended, Gram-positive bacilli, which have a characteristic growth on culture media.
Vaccinia. An accidental infection about the face may be mistaken for anthrax pustule. If inquiry into the attendant circumstances is not sufficient to exclude the graver disorder, a bacteriological examination should be made.

Primary syphilitic sore, if found on the face, is generally situated on the upper lip, although it may also occur upon an eyelid, the nose or elsewhere. It is not so indurated as when on the glans penis, but the surrounding oedema is more marked, and the neighbouring lymph nodes become enlarged. The condition is often missed because it is not expected. An absolute diagnosis can be made by finding the spirochaetes in the serum discharges from the ulcer, and by serological tests, although the latter may not yet be positive if the facial chancre is of recent date.

Insect bites or stings – from mosquitoes, gnats, bees, etc. – often cause large, lumpy, irritating swellings. The only difficulty in diagnosis is when the original bite or sting has become indistinguishable owing to infection with pyogenic organisms.

The various skin diseases that may be associated with swelling of the face are considered under BULLAE AND VESICLES (p. 76).

### FACE, ULCERATION OF

#### Barry Monk

Any persistent ulcerated lesion on the face, especially in a fair-skinned subject, raises the suspicion of malignancy and requires investigation, by biopsy if necessary, to determine the diagnosis (Box F.3). The commonest ulcerating skin cancer is the rodent ulcer (basal cell carcinoma) (Fig. F.12), and although this is traditionally associated with the elderly farmer with a lifetime of sun exposure, these lesions are not infrequently seen at a much younger age, and, thanks to recreational and holiday habits, in those with indoor occupations. Patients frequently associate the lesion with some minor incidental trauma (‘a scratch that just won’t heal up’) and this misleading history may delay the presentation and recognition of the true diagnosis. The hallmark of a rodent ulcer is its edge, which is raised and rolled with a pearly colour and crossed by multiple telangiectatic capillaries. Usually, there is central ulceration, but some lesions remain nodular or cystic for many months before ulcerating. Rodent ulcers can be deepy pigmented, making the true diagnosis less obvious, with simulation of a banal seborrhoeic wart or malignant melanoma. Squamous cell carcinoma is also common on sun-damaged skin, especially the lower lip, and may ulcerate.

#### Box F.3 Causes of face ulceration

<table>
<thead>
<tr>
<th>Benign</th>
<th>Infection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Excoriation</td>
<td>Syphilitic chancre</td>
</tr>
<tr>
<td>Trauma</td>
<td>Gumma</td>
</tr>
<tr>
<td>Artefacts</td>
<td>Yaws</td>
</tr>
<tr>
<td>Anaesthetic areas (trigeminal nerve ablation)</td>
<td>Leishmaniasis</td>
</tr>
<tr>
<td>Tumours</td>
<td>Lupus vulgaris</td>
</tr>
<tr>
<td>Basal cell carcinoma</td>
<td>Other</td>
</tr>
<tr>
<td>Squamous cell carcinoma</td>
<td>Pyoderma gangrenosum</td>
</tr>
<tr>
<td>Keratoacanthoma</td>
<td>Cancrum oris</td>
</tr>
<tr>
<td>Malignant melanoma</td>
<td>Dental sinus</td>
</tr>
</tbody>
</table>

Lesions begin as firm fleshy tumours, and grow slowly and asymmetrically. Keratoacanthoma is now regarded as a variant of squamous cell carcinoma; they present with the alarmingly rapid development of a smooth, symmetrical nodule with a central keratin plug, with is extruded leaving an ulcerating crater. If you are stoical enough not to excise the lesion, it commonly regresses spontaneously, leaving an irregular scar as a permanent reminder.
Ulceration may occur in malignant melanoma and is always a sinister sign, associated with a poor prognosis and the likelihood of early metastatic spread. Infectious causes of facial ulceration are also usually serious. Syphilis is no longer the rarity it was 20 years ago and is a diagnosis which must not be missed. A primary syphilitic chancre may occur anywhere on the face, but especially on the lips. It develops rapidly from a small nodule to an indurated, painless ulcer with associated marked lymphadenopathy. Its rapid growth should distinguish it from neoplasms, and Treponema pallidum can be found in large numbers on dark-field examination of the serous exudate. Serological tests will be positive 10–14 days from the onset of the chancre. A tertiary syphilitic gumma tends to ulcerate rapidly, extending at the margins and healing centrally.

Increasing population mobility means that cutaneous leishmaniasis is no longer merely seen in endemic areas. It must always be considered in the returning traveller, and may present with one or more facial ulcers. The lesion begins at the site of a sandfly (Phlebotomus) bite. Within a few weeks a livid papule develops and grows to 1–2 cm in size before ulcerating. The smear at this stage will be positive for Leishman–Donovan bodies. If untreated, most such ulcers heal with scarring in 12–18 months. In lupus vulgaris (cutaneous tuberculosis), which is now very rare, the ulceration is chronic. It begins with deep-seated nodules that, after a time, break down to form a granulomatous ulcer, covered with crusts. Around the edge, the characteristic ‘apple-jelly’ nodules may be seen. Necrosis of cartilage of the nose and pinna is not uncommon, but bone is never attacked (in contrast to syphilis and malignancy). Lupus vulgaris usually begins in childhood. Rare causes of facial ulceration include pyoderma gangrenosum, cancrum oris and ulcerating dental sinus (Fig. F.13).

Ulceration may be traumatic, for example in the habitual, and often subconscious, phenomenon of acne excoriée. Self-inflicted injury to the skin may be denied by the patient suffering from dermatitis artefacta, but the rather shallow history and the unnatural appearance of the lesions usually give a clue to the cause; there is invariably a history of psychological disturbance. Anaesthetic skin is soon traumatized and often prevented from healing by recurrent excoriation, and this can lead to extensive ulceration, as seen following surgery to the Gasserian ganglion for trigeminal neuralgia (Fig. F.14). A similar situation obtains in other causes of facial anaesthesia, including posterior inferior cerebellar artery thrombosis and syringobulbia.
Factitious disorders are conditions in which illnesses are simulated or induced, with deliberate deceit, but with no obvious motive or goal. Their prevalence is uncertain, but it is likely to be seriously underestimated as recognition is difficult unless the presentation is characteristic, persistent, repeated or closely observed – and suspected (Box F.4).

Broadly, there are three types of presentation: (i) the fabrication of symptoms or signs of illness; (ii) self-inflicted injuries to induce or simulate illness; and (iii) the aggravation or elaboration of established disease. Any physical or psychiatric condition can be feigned: even cases of feigned AIDS were reported within a year or two of its emergence. There seems to be no limitation to the ingenuity or self-inflicted suffering of some patients and, particularly among people with a medical background or training, the application of drugs, medical equipment or pathology samples to devise illness can be scarcely credible.

The first and greatest problem facing the doctor is to recognize that the presentation is not a bona fide illness. There are usually clues in the history, which may be dramatic but is more frequently vague, inconsistent and ‘hollow’, while pressing for better information can provoke resentment or anger. A detailed medical knowledge or excessive interest in the investigations or treatment may be evident. The rapid development of ‘complications’ or fresh, unrelated symptoms as the investigations undertaken begin to report negative results is most characteristic.

There are few readily identifiable presentations, but dermatitis artefacta is both common and may have a classically geometrical pattern to the lesions, with sharply demarcated edges; the surgical wound or ulcer that repeatedly breaks down is another suspicious scenario, with tampering sometimes apparent. Fever is frequently simulated usually by interference with the thermometer, but has been induced by injecting pyrogenic material. Bleeding from any orifice can be especially dramatic, although sometimes identifiable as self-inflicted upon careful examination – surreptitious anticoagulant ingestion or the addition of blood to samples can prove harder to identify, while the patient may occasionally induce anaemia before presenting. Gastrointestinal and endocrine disturbances are well documented, often involving drugs. Psychosis and bereavement are commonly feigned psychological presentations.

The diagnosis becomes even more difficult when the factitious behaviour is superimposed upon a genuine illness or abnormal test result, such as chest pain in the setting of electrocardiographic evidence of an old infarct, or when the patient accepts treatment knowing he should not have the treatment, such as with an established anaphylactic response to penicillin.

Finally, manufacturing factitious disorders in others is a rare but recognized phenomenon occurring very occasionally with health staff fabricating abnormalities in patients, and more often in parents inducing or simulating illness in their child or children. This form of the disorder, Munchausen’s syndrome by proxy, is associated with a factitious illness or somatization disorder in the mother, and differs from typical child abuse in that the parents usually make no attempt to disguise their children’s disorders or else give patently inadequate explanations; thus, the spectre of non-accidental injury is rarely first raised by the parents’ behaviour and attitudes. When a factitious condition is uncovered in an adult, it is always worthwhile reviewing the medical histories of any children (Table F.1). Munchausen’s syndrome by proxy should be investigated and managed from a forensic (evidence-gathering) perspective, and it represents a major child protection challenge. Therapy for the perpetrator is the secondary concern.

The differential diagnosis of factitious disorder is relatively straightforward compared with the problems usually encountered in establishing the nature of the condition.

### Table F.1 Differential diagnosis of factitious disorder

<table>
<thead>
<tr>
<th>Behaviour is evident and admitted</th>
<th>Behaviour is under voluntary control</th>
<th>Behaviour is directed towards an obvious goal</th>
</tr>
</thead>
<tbody>
<tr>
<td>Factitious disorder</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Deliberate self-harm</td>
<td>Usually</td>
<td>Yes</td>
</tr>
<tr>
<td>Hysteria</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Malingering</td>
<td>No</td>
<td>Yes</td>
</tr>
</tbody>
</table>

**Box F.4 Causes of factitious disorder**

**Most common**
- Personality disorder (antisocial, inadequate/dependent)
- Learning disability
- Brain damage
- Rare
- Schizophrenia
- Schizoaffective disorder
- Anorexia nervosa

**Less common**
- Depression
- Anxiety

Andrew Hodgkiss
Deliberate self-harm is also self-injurious behaviour, but it is overt and does not entail deception in the sense of intentionally mimicking other diseases. Conversion disorder involves functional complaints and signs, but these are not under conscious control and usually have identifiable gain (either primary gain, such as the resolution of psychological conflicts, or secondary gain, i.e. real-world gains such as being excused duties). Malingering is self-induced and consciously operated, but it is directed towards an understandable goal (such as cash benefits or avoidance of prison), which, once achieved, ends the behaviour.

The best known presentation of factitious disorder is Munchausen's syndrome. Typically, the patient is described as a young or middle-aged male, has repeated presentations and must wander. Their bizarre and multiple presentations, disruptive demanding behaviour, maladjusted unstable background and rootless existence are all hallmarks of severe personality disorder. It is because they are caricatures of the disorder that they are readily recognized, and their importance, numerically and in the literature, has undoubtedly become exaggerated.

In fact, the patient with factitious disorder is more likely to be female than male, to be undramatic, not to be admitted to hospital and to behave in a compliant manner. She tends to be a young adult, socially conforming, in employment and with an apparently stable family background. Pre-existing disease, ready access to drugs, and medical knowledge gained from training or working in healthcare or having a close relative in the health profession are all common. The personality is characterized as passive, dependent, sensitive, introverted, obsessional and with low self-esteem; sexual difficulties are common.

The condition sometimes represents a means of coping with an illness, a stress or a relationship problem in an individual who cannot articulate or assert, or it may be the expression of an underlying depressive illness. In acute cases, the outcome is favourable given appropriate management of the presentation and resolution of the primary problem. For those patients in whom the behaviour is repeated and arises in a more fundamental disorder of personality, themes of masochism and self-destruction figure persistently and prominently, self-injury is compelling and suicide a risk.

Finally, the doctor who establishes their patient is simulating illness is invariably faced with a key question – to confront or not to confront? The answer is almost always to confront but to approach the denouement gently, dispassionately, gradually and as part of a management plan, not in a hot-tempered, accusatory way. When confronted, some individuals with Munchausen's syndrome will get up, go and materialize elsewhere, but the remainder, the majority, will usually either respond positively to the suggestion of psychological help or will desist from their behaviour even if hotly denying the problem was self-induced. It helps if this process is carried out by the physician who assessed the patient and has made the diagnosis after careful evaluation; the psychiatrist is also then freed from implication in the detection process and can be seen as a helper after the event. Discovery can come as considerable relief for some patients who dislike their dishonesty as much as their doctor does, but no matter how skilfully and sensitively handled, these cases seldom prove satisfying and very rarely conclude with the doctor–patient relationship unscathed.

FAECES, COLOUR, CONTENTS AND MORPHOLOGY

Simon Anderson

Examining stool to determine consistency, shape, colour and content can provide some clue as to a patient's diagnosis, but findings are often inconsistent and usually can only be put into context after a diagnosis has been made.

STOOL CHARACTERISTICS

Stool characteristics are best described using the Bristol Stool Chart. To make a broad generalization, in the presence of diarrhoea, stools of large and small volume represent small- and large-bowel pathologies, respectively. Malabsorptive states (exocrine pancreas impairment: chronic pancreatitis, pancreatic cancer, cystic fibrosis; and, to a lesser degree, mucosal disease: coeliac disease, tropical sprue, lymphoma, lymphangiectasia, Whipple's disease, amyloidosis, systemic sclerosis) are associated with the passage of bulky, pale stools.

Hard- or pellet-like stools (Bristol type 1 and 2) are common in constipation. Ribbon-shaped stools are usually not diagnostic but occasionally are due to conditions causing narrowing of the distal colon (diverticular disease, tumour, Crohn's disease, anal stenosis).

The stools of bile salt malabsorption due to terminal ileal diseases such as Crohn's disease, lymphoma or infections (TB, Yersinia, HIV) are often watery. Cholera is associated with ‘rice-water’ stools and resulting dehydration.
**Stool pallor**, whether white or putty-coloured, is a sign of biliary obstruction resulting in decreased levels of stercobilinogen. Such conditions include common bile duct stone, ductal worms (ascarid or fasciola), haemobilia, cholangiocarcinoma, primary sclerosing cholangitis, ampullary adenoma/carcinoma, pancreatic head carcinoma, biliary atresia, surgical duct injury and anastomotic stricture. Steatorrhoeic stools are often described as being both pale and bulky.

Black bowel motions are frequently incorrectly labelled as melena. This term is reserved for altered blood that gives the stool a consistency of sticky black tar, with a characteristic pungent odour. Melena is usually a sign of upper gastrointestinal bleeding (peptic ulcer disease, oesophageal varices, reflux oesophagitis, gastritis, Dieulafoy’s lesion, cancer, angioectasia) but can also occur when bleeding originates from the distal small bowel or right colon (tumours, polyps, angioectasia, diverticula). Bleeding from the rest of the colon is usually described as maroon or burgundy, and bright-red (haematochezia) when from the distal colon (see below). Iron supplements will give stools a black appearance, but the stools may be solid and will have a slight dark-green tinge. Bismuth preparations and liquorice may blacken stools.

Poorly digested foods such as tomato skins or beetroot may be mistaken for blood, or during colonoscopy for being an adenomatous polyp.

Mucus, which is a normal constituent of stools, can be overproduced by large distal colonic polyps and is a common feature of irritable bowel syndrome (IBS). Mucus, when seen with blood is a feature of active ulcerative colitis. When whitish, strands of mucus can be mistaken for blood, or during colonoscopy for being an adenomatous polyp. The ova are oval with a clear, transparent shell detected on a direct faecal film or a slide mounted in saline or iodine solution, and they are commonly associated with iron-deficiency anaemia. Trichuris trichuria (whipworm) is very common, mainly in the tropics, and can present with diarrhoea, anaemia, pica and nocturnal pruritus ani. The worm is visible to the naked eye in the stool or perianal area.

**WORMS**

Several types of worm may appear in the stool: these include tapeworms, roundworms, threadworms and pinworms.

A common indication of tapeworm infestation is the passage per rectum of detached segments in either long or short tape-like strips. Close examination reveals the regular segmentation of a tapeworm, and examination with a lens reveals the glandular structure of the uterus in tapeworm segments. Patients may be symptomless or may complain of abdominal discomfort or diarrhoea; anaemia and eosinophilia may be present. The four forms of tapeworm which occur in the human intestine are Taenia solium (pork tapeworm), T. saginata (beef tapeworm), Hymenolepis nana (dwarf tapeworm) and Diphyllobothrium latum (fish tapeworm). T. saginata is the commonest tapeworm found in Britain. Microscopic examination of the faeces will show the characteristic eggs. Identification of the species is generally possible by the gravid proglottides. Tapeworms may also be seen on a straight X-ray film of the abdomen, or occasionally on barium meal examination.

The only roundworm which infests humans in Britain is Ascaris lumbricoides. Symptoms of intestinal colic or biliary obstruction, particularly in children, may occur together with pneumonitis, urticaria and eosinophilia. There may be no symptoms until a worm is found in the stool; typical ova may be discovered in faeces. They are of relatively large size and of oval shape. Enterobius vermicularis (known as threadworm in the UK and pinworm in the USA) is the most common helminthic infection in the US and Western Europe and generally occurs in large numbers. These can be detected by naked eye examination of the faeces. Each parasite is 3–10 mm in length and is colourless. They may be associated with frequency of micturition, pruritus ani, irritability and restlessness.

Other worm infections include hookworm (Ankylostoma duodenale and Necator americanus). The ova are oval with a clear, transparent shell detected on a direct faecal film or a slide mounted in saline or iodine solution, and they are commonly associated with iron-deficiency anaemia. Trichuris trichuria (whipworm) is very common, mainly in the tropics, and can present with diarrhoea, anaemia, pica and nocturnal pruritus ani. The worm is visible to the naked eye in the stool or perianal area.

**BLOOD IN STOOLS**

Black bowel motions are frequently incorrectly labelled as melena. This term is reserved for altered blood that gives the stool a consistency of sticky black tar, with a characteristic pungent odour. Melena is usually a sign of upper gastrointestinal bleeding (peptic ulcer disease, oesophageal varices, reflux oesophagitis, gastritis, Dieulafoy’s lesion, cancer, angioectasia) but it can also occur when bleeding originates from the distal small bowel or right colon (tumours, polyps, angioectasia, diverticula). Bleeding from the rest of the colon is usually described as maroon or burgundy, and bright-red (haematochezia) when from the distal colon. It is important to emphasize that there are many exceptions to this; for instance, heavy upper gastrointestinal bleeding can produce fresh rectal bleeding. Iron supplements will give stools a black appearance, but the stools may be solid and will have a slight dark-green tinge.
Bismuth preparations and liquorice also blacken stools. Fresh rectal bleeding (haematochezia) is often due to haemorrhoids (particularly when coating the outside of stools and when straining). Other causes include colorectal cancer, ulcerative colitis limited to the rectum (proctitis), polyps, rectal ulcer or anal fissure. Uncomplicated diverticular disease may give rise to heavy fresh rectal bleeding but never passage of intermittent small-volume blood unless it is accompanied by a degree of diverticular-associated colitis.

Colonic angiodysplasia (usually proximal colon) and rectal varices are less common causes of haematochezia. Colonic ischaemia, often of the splenic flexure, presents as abdominal pain and bloody diarrhoea in an elderly patient who may have suffered from recent hypotension (sepsis, major surgery). Intussusception is classically associated with the combination of abdominal pain and the passage of ‘redcurrant jelly’-like stools.

It is often suggested that blood mixed in with, or coating, the stool might be indicative of proximal and distal lesions, respectively. The identification of dark and bright red blood has similarly been used to suggest the site of pathology; however, none of these signs are diagnostic. Proximal colonic cancers may present with stools coated in bright red blood. Consequently, it is very important that all significant rectal bleeding is investigated with either colonoscopy or flexible sigmoidoscopy (depending on the age of the patient and nature of bleeding), as colorectal cancer is always a significant risk.

INVESTIGATIONS

The investigation of altered stools is guided by the above-mentioned features and their associations. The most accurate tests are those of colonoscopy and upper gastrointestinal endoscopy (gastroscopy). These procedures are superior to barium studies as they permit mucosal biopsy of the colon/terminal ileum and proximal small bowel (second part of the duodenum), respectively. CT colonography (CT enema, virtual colonoscopy) is accurate at detecting lesions ≥6 mm in size but biopsies cannot be obtained.

Blood tests will help to define associated nutritional deficiencies (ferritin, vitamin $B_12$ and serum/red cell folate) and, possibly, the underlying cause (anti-endomysial or tissue transglutaminase antibodies for coeliac disease, CA19.9 and CA125 for pancreatic cancer).

Five-day stool collection to assess fat content can define steatorrhoea, but this is rarely performed due to practical difficulties.

Tests to detect pancreatic function are seldom performed and usually only used for research. These include the bentiume test (BT-PABA), pancreolauryl test, intravenous secretin test and the Lundh test meal with duodenal aspirate for enzyme/bicarbonate measurement. The D-xylose absorption measures small bowel absorptive capacity.

Hydrogen/methane breath tests can detect lactose maldigestion and small bowel bacterial overgrowth.

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**FAECES, INCONTINENCE OF**

**Harold Ellis**

The individual who is affected by the inadvertent voiding of rectal contents per anum exists in a state of social alienation and professional isolation. Anorectal control is maintained under normal conditions by a combination of several factors, the most important of which include: (i) the internal anal sphincter; (ii) the external anal sphincter; (iii) the puborectalis muscle; and (iv) anorectal sensation. The role of the internal (involuntary) anal sphincter appears to be largely one of support, providing a ‘fine-tuning’ mechanism. Weakness of this muscle (e.g. following manual dilatation of the anus or sphincterotomy) leads to incontinence of flatus and soiling in the presence of diarrhoea, but not to major functional disturbance. The external (voluntary) anal sphincter can contract vigorously for approximately 60 seconds before fatiguing. This is probably a mechanism to prevent soiling (for a short period) should the anal sphincters become ‘threatened’ by the presence of loose stool in the rectum. A major contribution to anorectal control is provided by the contraction of the puborectalis muscle, which creates an angle between the lower rectum and upper anal canal (the anorectal angle). Sharp angulation permits a flap-valve mechanism to operate such that increases in intra-abdominal pressure cause the anterior rectal wall to close over the top of the anal canal, preventing rectal contents entering it. The sensation of a full rectum is probably caused by tension on pressure receptors situated in the pelvic floor rather than within the rectum itself. The discrimination of the nature of rectal contents is achieved by a simple locally mediated reflex whereby rectal distension (from flatus or faeces) initiates internal anal sphincter relaxation. A sample of rectal contents thereby intrudes into the anal canal and makes contact with the sensory-rich anoderm at the dentate line where it is perceived.

A complete classification of the causes of faecal incontinence is provided in Box F5. At the outset, it is of importance to establish the degree of disability,
Box F.5 Classification of the causes of faecal incontinence

<table>
<thead>
<tr>
<th>Normal sphincters and pelvic floor</th>
<th>Faecal impaction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Causes of diarrhoea (e.g. infection, inflammatory bowel disease)</td>
<td></td>
</tr>
<tr>
<td>Faecal fistula/colostomy</td>
<td></td>
</tr>
<tr>
<td>Abnormal sphincters and/or pelvic floor</td>
<td></td>
</tr>
<tr>
<td>Minor incontinence</td>
<td></td>
</tr>
<tr>
<td>Internal sphincter deficiency</td>
<td></td>
</tr>
<tr>
<td>– Previous surgery (e.g. anal dilatation, sphincterotomy)</td>
<td></td>
</tr>
<tr>
<td>– Rectal prolapse</td>
<td></td>
</tr>
<tr>
<td>– Third-degree haemorrhoids</td>
<td></td>
</tr>
<tr>
<td>– Idiopathic</td>
<td></td>
</tr>
<tr>
<td>Minor denervation of external sphincter and pelvic floor</td>
<td></td>
</tr>
<tr>
<td>Major incontinence</td>
<td></td>
</tr>
<tr>
<td>Congenital anomalies of the anorectum</td>
<td></td>
</tr>
<tr>
<td>Trauma</td>
<td></td>
</tr>
<tr>
<td>– Iatrogenic</td>
<td></td>
</tr>
<tr>
<td>– Obstetric</td>
<td></td>
</tr>
<tr>
<td>– Fractures of the pelvis</td>
<td></td>
</tr>
<tr>
<td>– Impalement</td>
<td></td>
</tr>
<tr>
<td>Denervation</td>
<td></td>
</tr>
<tr>
<td>– Obstetric</td>
<td></td>
</tr>
<tr>
<td>– Rectal prolapse</td>
<td></td>
</tr>
<tr>
<td>Peripheral neuropathy (e.g. diabetes mellitus)</td>
<td></td>
</tr>
<tr>
<td>– Cauda equina lesion (tumour or trauma)</td>
<td></td>
</tr>
<tr>
<td>– Tabes dorsalis</td>
<td></td>
</tr>
<tr>
<td>– Lumbar meningomyelocoele (spina bifida)</td>
<td></td>
</tr>
<tr>
<td>Upper motor neurone lesion</td>
<td></td>
</tr>
<tr>
<td>– Cerebral</td>
<td></td>
</tr>
<tr>
<td>– Multiple stroke</td>
<td></td>
</tr>
<tr>
<td>– Metastases and other tumours</td>
<td></td>
</tr>
<tr>
<td>– Trauma</td>
<td></td>
</tr>
<tr>
<td>– Dementia and other degenerative disorders</td>
<td></td>
</tr>
<tr>
<td>Spinal</td>
<td></td>
</tr>
<tr>
<td>– Multiple sclerosis</td>
<td></td>
</tr>
<tr>
<td>– Metastases and other tumours</td>
<td></td>
</tr>
<tr>
<td>– Degenerative diseases (e.g. vitamin B12 deficiency)</td>
<td></td>
</tr>
<tr>
<td>Rectal carcinoma</td>
<td></td>
</tr>
<tr>
<td>Anorectal infection (e.g. lymphogranuloma)</td>
<td></td>
</tr>
<tr>
<td>Drug intoxication (particularly in the elderly)</td>
<td></td>
</tr>
</tbody>
</table>

since the management of the patient with partial soiling secondary to a prolapsed haemorrhoid will clearly differ from that of a patient with frequent and incapacitating incontinence of formed stool. In all patients, a full clinical examination with special reference to the anorectum should be carried out. Digital examination of the anorectum will provide a subjective assessment of anorectal function, which ought to be supported, wherever possible: by (i) proctography; (ii) anal canal manometry; and (iii) electromyography of the external anal sphincter and puborectalis muscles.

**FAECAL INCONTINENCE IN THE PRESENCE OF NORMAL ANAL SPHINCTERS AND PELVIC FLOOR**

It is important to stress that the symptom of faecal incontinence need not necessarily imply deficiency of the anal sphincters or pelvic floor. Hence, any patient experiencing severe diarrhoea will frequently develop soiling of varying degree. The most common cause of faecal incontinence is therefore probably *gastroenteritis*. Patients with severe *inflammatory bowel disease* frequently state that the urgency and frank faecal incontinence is the most distressing aspect of the disease, and this, rather than the bleeding, may militate towards a surgical approach to management. Elderly patients and those who have *depressed cortical awareness* of rectal filling (e.g. following a cerebrovascular accident (CVA) or spinal cord section) may develop faecal impaction. Incontinence in these patients probably results from overactivation of a visceral reflex whereby the internal sphincter relaxes in response to rectal distension. A wide-open internal sphincter then permits the leakage of stool of looser consistency.

**MINOR FAECAL INCONTINENCE**

This is defined as the inadvertent loss of flatus or liquid stool per anum, and is usually the consequence of a weak internal anal sphincter. This situation may arise secondary to some surgical procedures (e.g. manual dilatation of the anus) or be caused by a stretch effect in patients with a full-thickness *rectal prolapse* (Fig. F.15) or with *third-degree haemorrhoids* (Fig. F.16). In some patients, internal sphincter dysfunction is observed *without any underlying cause* being apparent; in these patients, there may be disease affecting the autonomic supply to this muscle. Finally, minor degrees of incontinence may result from *denervation* and other *injuries* affecting the external anal sphincter and pelvic floor; these are discussed below.

**MAJOR FAECAL INCONTINENCE**

This is defined as the inadvertent and frequent loss of fully formed stool per rectum and, as such, represents the most severe degree of functional impairment of the anorectum. *Congenital abnormalities* of the lower gut may be associated with anorectal incontinence, particularly in some forms of rectal atresia where there

**Figure F.15 Complete rectal prolapse.**
has been a total failure of development of the pelvic floor musculature. Traumatic damage may be inflicted on the external sphincter during vaginal delivery, in which case the damage sustained is usually confined to the anterior section of the sphincter (third-degree perineal tear) or by the surgeon during the treatment of anal fistula when perhaps the puborectalis muscle or too much external sphincter muscle has been inappropriately divided. The pelvic floor can be damaged by ‘shearing’ forces when there has been complete disruption of the bony pelvis following compression injury to the pelvis. Rarely, impalement injuries to the anal sphincters and pelvic floor can lead to severe functional loss.

The greatest number of patients presenting for treatment of major faecal incontinence are found to have denervation of the striated component of the anal sphincter musculature. The source of nerve damage seems to be local (i.e. pudendal) in the majority, and a major factor would appear to be traumatic childbirth in which the nerves are subjected to undue compression and stretching forces. Damage may also be sustained in patients who strain excessively with defecation, and less commonly in patients with peripheral neuropathies, particularly diabetes mellitus.

Finally, very rarely, lower motor neurone lesions can be the consequence of cauda equina tumours. If there is a history of severe perineal pain and the history of incontinence is brief, this diagnosis should be considered and a spinal magnetic resonance image obtained.

Upper motor neurone lesions cause faecal incontinence for imprecise reasons. There is little doubt that interruption of suprasegmental control causes incontinence, partly as a consequence of a motor deficit and partly because of sensory loss, which in turn leads to impaction.

Rarely, rectal carcinoma and infection (specifically by lymphogranuloma venereum) can give rise to extensive destruction of the pelvic floor such that faecal incontinence might develop.

FAINTS
Mark Kinirons

(See also FITS AND CONVULSIONS, p. 205; VERTIGO, p. 736.)

Attacks of transient loss of consciousness are common, and the causes range from simple vasovagal attacks to epilepsy. Furthermore, a complaint of ‘fainting’ may not always imply actual loss of consciousness; some patients may mean no more than a feeling of unsteadiness or ‘light-headedness’. It is important, as always, to obtain from the patient and, whenever possible, from a witness a precise description of the nature of the attacks. The term ‘syncope’ has a more exact connotation, and can be defined as ‘transient loss of consciousness due to a reduction in the cerebral blood flow’. It is this condition with which this discussion will be mainly concerned.

The common causes of syncope are summarized in Box F6.

VASOMOTOR SYNCOPE
This rather imprecise term will be taken to include the large group of conditions in which the cardiac output and blood pressure fall as a result of a

<table>
<thead>
<tr>
<th>Vasomotor</th>
<th>Cardiac syncope</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Vasovagal attacks</td>
<td>• Stokes–Adams attacks in atrioventricular block</td>
</tr>
<tr>
<td>• Postural hypotension</td>
<td>• Paroxysmal dysrhythmias</td>
</tr>
<tr>
<td>• Carotid sinus syncope</td>
<td></td>
</tr>
</tbody>
</table>

<table>
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<tr>
<th> </th>
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</tr>
</thead>
<tbody>
<tr>
<td>• Central circulatory obstruction (e.g. aortic stenosis)</td>
<td>• Cerebrovascular disease</td>
</tr>
<tr>
<td>• Cyanotic attacks in congenital heart disease</td>
<td></td>
</tr>
</tbody>
</table>
sudden fall in peripheral resistance and in the central venous pressure.

**Vasovagal attacks**

These are extremely common and are almost always of no serious significance. Many predisposing factors are known, including emotion, fatigue, prolonged standing and chronic illness of almost any kind. There are other, more serious, conditions of which an apparently simple faint may be a manifestation. Haemorrhage causes syncope as a direct result of the fall in central venous pressure. If the bleeding is external, no diagnostic difficulty arises, but with internal bleeding (e.g. into the gastrointestinal tract), syncope may occur before there is any direct evidence such as haematemesis or melaena. The mechanism whereby haemorrhage causes syncope can also operate in a few patients with very large varicose veins or angiomatous malformations in the legs in which the blood accumulates in the upright posture. Severe pain can also cause syncope, as in a dissecting aneurysm or myocardial infarction, although, in the latter condition, there may also be a fall in cardiac output sufficient on its own to cause syncope. In elderly patients especially, syncope is quite commonly the presenting symptom of myocardial infarction; on recovering consciousness, the patient may not complain of chest pain, being perhaps more preoccupied with any trauma they may have suffered in their fall.

The clinical picture is well known – not least to the many medical students who have fainted at their first operation. The patient is nearly always in the upright position; indeed, a ‘faint’ occurring in the recumbent position is good evidence of some cause other than a vasovagal attack. A minor exception to this rule is the syncope experienced by some pregnant women while lying on their backs; this is probably due to pressure by the uterus on the inferior vena cava producing a fall in venous return. Prodromal symptoms include a feeling of weakness, nausea, sweating and epigastric discomfort; within a few seconds or minutes, the patient falls unconscious. The pulse is of small volume and slow, and the blood pressure is very low; the face is pale, and the skin cold and sweating. Incontinence of urine and muscular twitching are rare, but they can occur in transient loss of consciousness from any cause and do not necessarily imply that the attack is epileptic. Recovery is rapid as the cerebral blood flow increases in the recumbent posture, unless the patient is prevented from falling as in a crowd or by well-meaning bystanders. Weakness and nausea may persist for some time after recovery of consciousness.

The term ‘vasovagal’, which has been used to denote this type of vasomotor syncope, is now widely accepted. However, in the past it was used, originally by Sir William Gowers, to describe curious ‘seizures’ that resembled the attacks described above in most respects. They differed only in that loss of consciousness was rare and no precipitating cause could usually be found. A theory that they represented a specific entity and were possibly epileptic in nature has now been discredited, and it seems certain that these attacks were no more than a mild form of vasomotor syncope occurring in unusually susceptible subjects. All practising physicians have seen patients who faint not only from venepuncture but from a tourniquet or sphygmomanometer cuff being put around an arm.

**Postural hypotension**

This condition overlaps with vasovagal syncope, which, as has been said, almost always occurs when the patient is upright. It is, however, worth distinguishing a group of patients who have a steep fall in blood pressure whenever they stand upright. In some of these patients, a neurological cause for the failure of vasoconstriction and other compensatory mechanisms can be identified. To others, the label ‘idiopathic’ has been applied but, in many of these, it is probable that detailed investigation would localize a neurological lesion.

Many drugs can cause a marked postural fall in blood pressure. Among these are hypotensive agents such as guanethidine and bethanidine; nitrates, phenothiazine derivatives, monoamine oxidase inhibitors, imipramine, barbiturates, amitriptyline and other psychotherapeutic agents have also been incriminated. The hypotensive agent prazosin seems to be unusually liable to cause sudden loss of consciousness for periods of time ranging up to 1 hour. It is not known whether this is always due to postural hypotension, and it may be a specific side effect of this drug. Lesions of peripheral nervous pathways can produce a similar effect by interruption of the afferent or efferent pathways of reflex arcs. Thus, postural hypotension is well known in tabes, diabetes and acute polyneuritis, and has been described in alcoholic and carcinomatous neuropathy and in porphyria. Lesions of the central pathways are less easy to demonstrate, but degeneration of the intermediolateral column in the spinal cord, vascular lesions of the brainstem and craniohypophyngioma, and other parasellar tumours, possibly involving the hypothalamus, have been demonstrated in some cases. Many of these neurological conditions have in common an abnormal
response to the Valsalva manoeuvre in that the blood pressure continues to fall throughout the period of strain and no overshoot or reflex bradycardia occurs.

**Cardiot sinus syncope**

In many subjects, massage of a carotid sinus can cause bradycardia with some fall in blood pressure. In some, mostly elderly patients, these changes may be more marked, and this increased sensitivity of the carotid sinus reflex can be produced by neoplastic or inflammatory lesions in the neck or by digitalis intoxication. In a few patients, the haemodynamic changes may be so profound and the reflexes so easily elicited, as by a tight collar, shaving or turning the head, that recurrent syncope occurs. Various types of carotid sinus syncope have been described with or without bradycardia in addition to the hypotension, but the distinction is largely of academic interest. The carotid sinus is innervated by the glossoopharyngeal nerve, and a rare condition that may be related to carotid sinus syncope is the fainting sometimes associated with glossoopharyngeal neuralgia, a condition similar to trigeminal neuralgia but causing pain in the tongue, pharynx and ear.

**Cardiac Syncope**

In this group of conditions, the cardiac output falls as a result of a primary cardiac lesion. It differs from vasomotor syncope in that attacks are much less closely related to the upright posture.

**Stokes–Adams attacks**

These are due to cardiac arrest, usually in asystole but occasionally in ventricular fibrillation, on a basis of atrioventricular block. Loss of consciousness can occur in any posture, and abrupt. The patient is pale and pulseless; respiration continues. After about 15–20 seconds, twitching may begin due to cerebral anoxia. The attack usually lasts for about 30 seconds, but it may last longer and death may result. On recovery, the patient becomes flushed; this is due to well-oxygenated blood that has been in the pulmonary capillaries during the period of circulatory arrest being flung into systemic capillaries that are widely dilated as a result of the accumulation of vasodilator metabolites. Occasionally, if the attacks occur when the patient is asleep, the only complaint may be of waking with the face feeling hot and flushed.

**Paroxysmal dysrhythmias**

A paroxysm of tachycardia with a heart rate much in excess of 200 beats per minute may cause syncope as diastolic filling of the heart is markedly reduced at these heart rates.

**Central circulatory obstruction**

Syncope on effort and (more seriously) at rest is a well-recognized feature of aortic stenosis. The mechanism is not clear as this valve lesion is not associated with a low cardiac output unless failure has occurred. It may be that baroceptors within the left ventricular wall, stimulated by the very high pressure, are in some way responsible. Effort syncope is also not uncommon in other obstructive lesions such as pulmonary stenosis and severe pulmonary hypertension, but it is rare in mitral stenosis. The acute circulatory obstruction produced by massive pulmonary embolism or by the impaction of a left atrial thrombus or myxoma in the mitral orifice may also cause syncope. Obstruction to cardiac filling due to cardiac tamponade and constrictive pericarditis can have the same effect.

**Cyanotic attacks**

Syncope in Fallot’s tetralogy and other types of cyanotic congenital heart disease is due not so much to a fall in cardiac output as to a sudden increase in the vena-arterial shunt. This can be due to a fall in systemic resistance or to an increase in the severity of a muscular infundibular stenosis. There is little change in blood pressure, but the patient becomes deeply cyanosed, and the murmur of pulmonary stenosis becomes much softer as the greater part of the systemic venous return is shunted into the aorta via the ventricular septal defect.

**Ischaemic heart disease**

Syncope due to acute myocardial infarction has already been discussed under ‘Vasomotor syncope’, above.

**Syncope due to Cerebral Vascular Disease**

Fainting is rare as a symptom of disease of the carotid arteries and their branches, but it is more common with stenosis or occlusion of the vertebral arterial system. Atherosclerosis of the vertebral or basilar arteries and external compression of the vertebral arteries by cervical spondylsis, or in the Klippel–Feil syndrome, can be responsible for syncopal attacks that may be induced by sudden rotatory movements of the neck.

**Miscellaneous**

There is a group of conditions in which syncope is associated with a rise in intrathoracic and intra-abdominal pressure. This includes cough syncope, in which loss of consciousness occurs at the end of a violent paroxysm of coughing. A series of coughs produces the same circulatory effects as a Valsalva manoeuvre, and a marked fall in cardiac output and blood pressure – probably combined with some degree
of arterial hypoxaemia – is the mechanism of the syncope. The *breath-holding attacks* of early childhood producing syncope and cyanosis probably have a similar mechanism. The mechanism of *micturition syncope* is not fully understood. It occurs typically when, after heavy beer-drinking, the subject rises in the middle of the night to pass urine. The sudden assumption of the upright posture and the vasodilatory action of alcohol are certainly relevant factors, and this condition may be nothing more than a vasomotor syncope. However, it is possible that afferent impulses from the bladder may play some part.

**DIFFERENTIAL DIAGNOSIS**

The most important condition from which syncope must be distinguished is *epilepsy*. This can be very difficult, but a careful history will usually resolve the problem. Epileptic attacks are characteristically stereotyped in their nature and duration, and often occur without warning. Even if there is an aura, it bears little resemblance to the prodromal symptoms of syncope except, possibly, in the case of complex partial seizures. Most varieties of syncope – with the notable exception of cardiac syncope – occur almost exclusively in the upright posture, whereas the onset of an epileptic fit is unrelated to posture. The typical tonic and clonic phases of major epilepsy should not be confused with the minor twitches that may occur in syncope if a history can be obtained from a reliable eye-witness. Urinary incontinence, a very common feature of epilepsy, is not a major differentiating factor as it can occur in severe or prolonged syncope. Electroencephalography is certainly of value in some cases, but even with this aid some doubt may remain.

Consciousness is not lost in *vertigo*, so that a history from the patient him- or herself should elicit an account of the typical sensation of rotation. However, the use by the patient of such terms as ‘giddiness’ to describe the premonitory symptoms of vasomotor syncope, together with the nausea that is common to both conditions, may confuse the unwary. *Hysterical attacks* are nearly always described by the patients as ‘faints’, but the gracefully dramatic fall into a convenient armchair bears no resemblance to syncope in which the patient collapses like a house of cards. Swoons are out of fashion, unless one includes the hysterical faints of teenage girls at pop concerts. It is possible that the so-called ‘Toronto blessing’ experienced in some charismatic church services is of the same nature.

*Hyperventilation* is usually a hysterical phenomenon, and the feeling of light-headedness induced may be described as faintness. Here, as always, a careful history will resolve the issue, but it is worth recalling that hyperventilation followed by a Valsalva manoeuvre will infallibly cause loss of consciousness. This is known variously as the ‘mess trick’ or the ‘fainting lark’.

**FALLS**

Mark Kinirons

Elderly people are liable to fall for many reasons other than ‘drop attacks’, and causes such as postural hypotension, lack of concentration, the effects of sedating medicines, and disturbances of vision, tripping and postural instability must be excluded from the diagnosis.

Falls are common. They increase with age and dependency (being most frequent in nursing homes). There are many risk factors for falls:

- Muscle weakness
- History of falls
- Gait deficit
- Balance deficit
- Use of assistive device
- Visual deficit
- Arthritis
- Depression
- Cognitive impairment
- Age over 80 years

In addition, falls have many causes:

- Accident/environment: 31 per cent
- Gait/balance/weakness: 17 per cent
- Dizzy/vertigo: 13 per cent
- Drop attack: 9 per cent
- Confusion – acute/chronic: 5 per cent
- Postural blood pressure: 3 per cent
- Visual disturbance: 2 per cent
- Syncope: 0.3 per cent
- Other: 15 per cent
- Unknown: 5 per cent

It is essential to make an accurate diagnosis, and a diagnosis can indeed be made in up to 95 per cent of all falls. This allows treatments to prevent further falls. In addition, consideration should also be given to preventing fractures due to falls. This includes treatment for osteoporosis.

**FASCICULATION**

Mark Kinirons

Fasciculation is one of the classical features of a lower motor neurone lesion. It is observed clinically as an intermittent twitching movement due to the
contraction of groups of muscle fibres supplied by the affected nerve. The term ‘fibrillation’ is applied to the contraction of single muscle fibres. Fibrillation is recorded electrically rather than observed clinically. Fasciculation can occur when only part of the lower motor nerve is affected.

Although fasciculation can occur with inflammatory or compressive lesions of peripheral nerves, it is in chronic degeneration of anterior horn cells that it is most conspicuous. It is thus a common and an important diagnostic feature of motor neurone disease when the lower motor nerve is affected. Intermittent flickering movements, often around the shoulder girdle or in the thenar muscles, and sometimes felt by the patient, may be an early sign of the disease. Inspection over 2 or 3 minutes may be necessary before the tell-tale movement is detected. Sometimes it can be provoked by a light tap over the muscles with a tendon hammer or flick of the finger.

It is important to note that fasciculation can also be benign. Transitory twitching of the facial muscles, particularly the orbicularis oculi, is common experience in normal people; so too is fasciculation in calf muscles, sometimes after unaccustomed exercise. Such benign fasciculation, particularly when it is prolonged or extensive, as it occasionally is, can be a cause of major anxiety in doctors and others unfamiliar with the features of motor neurone disease.

FATIGUE

Mark Kinirons

Fatigue is a normal reaction to exertion, but it may be produced excessively by minor exertion in any condition of ill-health, whether it be organic, psychogenic, or both. Fatigue may be caused by malignant disease, infections and their sequelae, anaemia and states of low cardiac output, endocrine and metabolic disorders and malnutrition. It may be due to lack of sleep, boredom, stress or overwork. The so-called ‘combat fatigue’ seen in the troops in the Second World War resulted more from the mental than the physical stress of active service. Fatigue on waking in the morning is more commonly due to mental rather than physical factors, but it may be due to either or both. Depending on the cause, fatigue may be constant or episodic and in some cases, as in myasthenia gravis, may be anticipated so that physical effort is avoided to prevent the later onset of fatigue.

Fatigue is at least as often psychological in origin as due to organic disease, but many cases are multifactorial in origin. The following list, although inevitably incomplete, includes most of the common causes.

EMOTIONAL AND PSYCHOLOGICAL CAUSES

These include continued unhappiness, boredom, disappointment, overwork, lack of sleep, anxiety and depression. Neuasthenia, a nineteenth-century diagnosis, is undergoing something of a revival.

MALIGNANT DISEASE

Any neoplastic disease can be associated with fatigue, which may be a very early symptom – sometimes earlier than loss of weight.

CHRONIC INFECTION

Tuberculosis, brucellosis, infective endocarditis and toxoplasmosis are examples of conditions of which the most obvious manifestation may be fatigue.

HUMAN IMMUNODEFICIENCY VIRUS

Fatigue is a significant problem for patients suffering from HIV disease, and is often under-reported by patients and therefore undertreated by physicians. It tends to be unpredictable and cyclical in nature, although there are some who suffer unrelenting fatigue. The degree of fatigue can be quite debilitating, interfering considerably with requirements for daily living and, without a patient carer, leads to an inability to wash, shop, eat and comply fully with complex medical regimens. The fatigue is often explained by just having the HIV disease itself, although it can be quite severe in patients who do not have a significant viral load. There are specific conditions that may make the fatigue worse, and these should be sought out since therapeutic intervention may be of benefit. Hypogonadism associated with decreased testosterone level, suprarenal insufficiency, methaemoglobinemia secondary to dapsone treatment, anaemia, depression and malnutrition are among the associated conditions contributing to fatigue, and should all respond to therapy. Appropriate exercise programmes have also been shown to be of benefit in improving energy levels.

POST-VIRAL FATIGUE SYNDROME

Many viral infections, of which infectious mononucleosis is the best-known example, may be followed, sometimes for many months, by profound fatigue; this may be curiously episodic. The status of myalgic encephalomyelitis (ME)/chronic fatigue syndrome as a specific entity is unclear. Some believe that it is a severe variety of the post-viral syndrome. Others maintain that it is a different condition. It has been suggested that graded exercise and cognitive behavioural therapy are beneficial in many cases of the post-viral syndrome and ME.
TISSUE HYPOXIA
Anaemia of any type is a common cause of fatigue, as are conditions in which the cardiac output is chronically low. These include cardiac failure, severe pulmonary hypertension, valvular heart disease (e.g. aortic and mitral stenosis), Addison’s disease and excessive diuretic therapy.

CONNECTIVE TISSUE DISEASES
These include rheumatoid arthritis, systemic lupus erythematosus, polyarteritis nodosa, polymyalgia rheumatica, giant-cell arteritis and polymyositis.

ENDOCRINE AND METABOLIC DISORDERS
Apart from Addison’s disease (see above), many conditions in this category, such as hypothyroidism, mast cell disorders, renal or hepatic failure and diabetes mellitus, can present with fatigue.

MALNUTRITION
Whether this is due to dietary deficiency or to conditions associated with chronic diarrhoea such as coeliac disease, ulcerative colitis or Crohn’s disease, fatigue is likely to be present.

CHRONIC PAIN
Fatigue can be due to persistent chronic pain causing discomfort by day, as in osteoarthritis, or lack of sleep at night, as in some cases of Paget’s disease or metastatic disease of bone.

MUSCULAR WEAKNESS
Chronic neurological disorders, such as multiple sclerosis, motor neurone disease and especially myasthenia gravis, in addition to the myopathies, are typically associated with excessive fatiguability.

CHRONIC DRUG INTOXICATION
The long-term administration of beta-adrenergic blocking agents is an important cause of fatigue. Alcohol abuse and the chronic administration of benzodiazepines have a similar, but less specific, effect.

DRUG WITHDRAWAL SYNDROMES
Fatigue may be a consequence of the withdrawal of addictive drugs such as morphine and diamorphine. It can also occur after the withdrawal of corticosteroids, benzodiazepines, alcohol and antidepressants.

FINGERS, CLUBBED
Alex West
The four features of finger clubbing are: loss of the nail bed angle; increased nail curvature (Fig. F.17); fluctuation of the nail bed; and drumstick-like swelling of the terminal phalanx. Identification is easy when all four features are present, but dispute is common when only some occur. The first two criteria must be present for finger clubbing to be diagnosed; thus, nail bed fluctuation on its own does not constitute clubbing. The first two criteria combine to increase the hyponychial angle, subtended at the nail base by the skin crease at the dorsum of the distal interphalangeal joint and the skin immediately below the free edge of the nail (Fig. F.18). This angle, which is best measured from a shadowgram, is usually less than 190° in normal subjects, but exceeds 195° in individuals with clubbing. Clubbing may affect the toes as well as the fingers but is usually more obvious in the hands. Clubbing is usually bilateral, but occasionally it may be unilateral in the presence of local vascular abnormalities, such as a subclavian aneurysm, an arteriovenous fistula or disruption of the cervical sympathetic nerves. The pathophysiology of finger clubbing is not understood.

Figure F.17 Clubbing of fingers of both hands due to bronchiectasis.

Figure F.18 Measurement of the hyponychial angle.
Finger clubbing is an important physical sign, and may be the first and only sign of serious organic disease. The relative frequencies of causes will vary between countries; typical causes in the UK are listed in Box F.7.

Among pulmonary diseases, lung carcinoma is the most common cause of finger clubbing, and it is wise always to consider this diagnosis. Other intrathoracic malignancies, including mesothelioma and lymphomas (and rarely secondary carcinomas or sarcomas), can cause finger clubbing. Intrapulmonary sepsis is associated with clubbing, although this is now an unusual cause in Britain. Chronic bronchitis and emphysema do not cause clubbing. Diffuse pulmonary fibrosis may be associated with clubbing, but different causes vary markedly in the frequency of this association. Clubbing is common in idiopathic pulmonary fibrosis, but rare in extrinsic allergic alveolitis and sarcoidosis. Pulmonary asbestosis is the only pneumoconiosis directly associated with clubbing.

Among diseases of the cardiovascular system, cyanotic congenital heart disease is almost always associated with finger clubbing in those surviving beyond infancy. However, clubbing does not occur in non-cyanotic congenital heart disease such as uncomplicated atrial or ventricular septal defects or persistent ductus arteriosus. Clubbing is a well-recognized, but rather uncommon, feature of infective endocarditis; its absence must not be regarded as important evidence against the diagnosis. It usually appears 6 weeks or more after the onset of the illness, but can occasionally develop within a month. Mild finger clubbing develops late in some cases of cirrhosis of the liver, especially in biliary cirrhosis, in coeliac disease and, rarely, in ulcerative colitis and Crohn’s disease.

Clubbing due to any cause may progress to hypertrophic pulmonary osteoarthropathy, although this is almost always due to lung carcinoma. Periosteal new bone formation will then be evident, especially at the distal ends of long bones at the affected joints. Familial forms of clubbing and of hypertrophic osteoarthropathy are rare. The presence in family members and the absence of associated disease provide clues to the diagnosis. Hereditary osteoarthropathy is associated with thickening of the skin of the hands and face, the latter giving rise to a characteristic appearance of large features with coarse, deeply creased skin. This rare familial condition is called ‘pachydermoperiostosis’.

Patients are rarely aware of their clubbed fingers. Thus, statements that there has been no recent change in their fingers should not be taken to indicate congenital clubbing.

FINGERS, DEAD (WHITE, COLD)

Melvin Lobo

RAYNAUD’S PHENOMENON

The digital arteries of the fingers serve two purposes: (i) they supply blood for nutrition of the finger; and (ii) by controlling blood flow through the skin of the fingers, they vary the heat loss and assist in regulating the core temperature of the body. Thus, on exposure of the body to cold, these arteries normally constrict, reducing blood flow through – and heat loss from – the fingers. In subjects with Raynaud’s phenomenon, this reflex appears to be excessive and the digital arteries open up again and blood flushes vigorously through the fingers, often causing throbbing discomfort (‘rewarming pain’). During severe episodes, pallor of the digits may be followed by cyanosis and numbness prior to rewarming. In such instances, the classical progression of triphasic digital colour changes is observed: white-blue-red. Other sites including the toes, tongue, nose, ears and nipples can also be affected.

Raynaud’s phenomenon occurs on a worldwide basis, although it is more prevalent in cold climates. It occurs
in about 3–5 per cent of healthy young females in Britain, and is far less common in males. Primary Raynaud’s phenomenon occurs in the absence of underlying disease, whereas secondary Raynaud’s phenomenon is associated with numerous disorders.

Primary Raynaud’s phenomenon

Primary Raynaud’s phenomenon has an earlier age of onset than secondary – typically between the ages of 10 and 30 years – with a milder disease course. The nailfold capillaries and fingers remain healthy and normal in appearance, and digital ulcers or gangrene rarely develop. In addition to these clinical features, a normal erythrocyte sedimentation rate and a negative test for antinuclear antibody help to distinguish it from secondary Raynaud’s. A family history is common and noted in up to 25 per cent of first-degree relatives of patients with primary Raynaud’s phenomenon. It is important to distinguish primary Raynaud’s phenomenon with predominantly unilateral symptoms from vascular thoracic outlet syndrome in which the doctor can induce symptoms of pallor and diminished pulses through physical manoeuvres.

Secondary Raynaud’s phenomenon

Much less common is Raynaud’s phenomenon secondary to some underlying disease. In these patients, the phenomenon usually appears later in life – often in middle age – although it can occur in younger patients. Men are affected more often than women in primary Raynaud’s phenomenon. From the beginning, the digital ischaemia tends to be more severe and is often at first asymmetrical. Recurrent and prolonged digital ischaemia eventually causes the fingers to change in appearance, becoming shrunk with tight skin and loss of subcutaneous tissue. Ulcers commonly appear under the fingernails (Fig. F.19) and, when healed, leave puckered scars. Repeated attacks lead to loss of tissue of the terminal phalanx with resorption of the phalanx and curved overhanging nails. Causes of secondary Raynaud’s phenomenon are listed in Box F.8.

Scleroderma is the most common cause of secondary Raynaud’s phenomenon, occurring as the presenting complaint in 70 per cent of cases. It is usually the variety of scleroderma known as the CRST syndrome (calcinosis, Raynaud’s phenomenon, sclerodactyly and telangiectases). Initially, only one or two of these four components of the syndrome will be present – usually the Raynaud’s and the telangiectases. The telangiectases are first seen on the nail bed, but later larger ones appear on the fingers and face. Subcutaneous calcification eventually appears, which is at first felt as tender nodules under the skin of the fingers, but eventually extrudes through the skin (Fig. F.20). Although severe involvement of the fingers often leads to a loss of digits, it does not usually affect vital organs and does not normally shorten life expectancy. Progressive systemic sclerosis with widespread skin involvement and involvement of vital organs is a rare cause of Raynaud’s phenomenon, and ulceration and gangrene of digits is unusual.

Systemic lupus erythematosus (SLE) may cause severe digital arthritis with Raynaud’s phenomenon and repeated attacks of digital gangrene causing extensive loss of digits. Rheumatoid arthritis can similarly affect the fingers with digital gangrene. Vibration injury, as seen in foundry workers using pneumatic-powered, hand-held buffers and grinders, caulkers and welders in the ship-building industry and forestry workers...
using hand-held, power-driven saws, can cause severe Raynaud’s phenomenon. The disorder may appear within a few months in foundry workers, but takes longer to appear in shipyard and forestry workers. In addition to the white fingers, these patients develop numbness and tingling of the fingers. Although the Raynaud’s phenomenon can be very severe and a considerable nuisance during cold weather, ulcers and gangrene generally do not occur.

Beta-blocking drugs used in the treatment of angina and hypertension commonly cause cold hands, but they do not seem to induce classic Raynaud’s phenomenon (i.e. digital vasoconstriction) in patients who would not otherwise have Raynaud’s. However, these drugs may make Raynaud’s worse in those already suffering from the condition.

**PERSISTENT DIGITAL ISCHAEMIA**

Sudden onset of ischaemia of one or more digits persisting for days, weeks or months (persistent digital ischaemia) is not uncommon in the middle-aged and elderly. On examination, the finger is usually blue and cold, but capillary circulation is present and the finger usually survives. In younger patients, this condition is usually due to some form of arteritis (e.g. SLE or rheumatoid disease), but in older patients investigation usually fails to reveal any abnormality except for the presence of atheroma. In these patients, the ischaemia is due to rupture of an atheromatous plaque higher in the arterial tree with plaque debris embolizing the digit (Fig. F.21). Digital ischaemia may also be due to more distant emboli (e.g. from the heart in patients with atrial fibrillation, or via a patent foramen ovale in patients with thrombophilia and venous thrombi; in these cases the ischaemia is often more significant and widespread.

**FROSTBITE**

Apart from provoking Raynaud’s phenomenon in predisposed individuals, prolonged exposure of the fingers to cold (e.g. in hill-walkers or outdoor workers in winter) may result in the patient complaining of cold, dead, numb fingers due to freezing of the superficial layers of the skin. In the early stages, there are white and dead patches of skin on the fingers, but later and in more severe cases, gangrene of the skin appears and may envelop the whole digit. Although it appears alarming at first, the gangrene is limited in most cases to the superficial layers of the skin, and the skin will eventually peel off, leaving a normal digit beneath.

**FINGERS, SKIN AFFECTIONS OF**

Barry Monk

The skin of the fingers is particularly prone to those dermatoses which are influenced by environmental exposure, such as cold, ionizing radiation and chemicals. The fingers are often the site of inoculation of infectious skin conditions. A list of conditions commonly seen on fingers, divided into morphological types, is provided in Box F.9. The conditions, listed by anatomical site, are detailed in Box F.10. Some examples are shown in Figs F.22–F.25.

Viral warts, often multiple and frequently periungual, are commonplace and readily identifiable; in older subjects, however, the diagnosis of a solitary viral wart must be treated with some circumspection, as it may mimic a squamous cell carcinoma or a small plaque of Bowen’s disease (squamous cell carcinoma in situ).

Orf, presenting in a rather dramatic fashion as a rapidly developing solitary inflamed blistering nodule on the
finger, is a viral infection acquired from lambs, and is commonly seen in patients in rural areas in the spring time. It spontaneously regresses in 2 to 3 weeks. Farmers, aware of the condition, rarely seek medical assistance, but visitors from urban areas may return home, having forgotten the contact with animals, and puzzle their doctor!

One condition peculiar to the finger is that of paronychia, an acute or chronic infection of the nail fold. The most important precipitating factor is loss of the ‘seal’ at the nail quick. It is commonly seen in those who have their hands immersed in water.
for long periods, such as barmaids and cleaners. *Candida albicans, Pseudomonas* and *Staphylococcus aureus* can all be causes, but the condition is difficult to eliminate unless the hands are kept scrupulously dry long enough for the natural seal to re-form. Acute paronychia can also be caused by *herpes simplex* (herpetic whitlow), and the presence of a painless ulcer at the lateral margin of the nail fold, frequently of the index finger, should arouse suspicion of a *syphilitic chancre*. Positive serological tests after the 10th to 14th day, or the development of a secondary rash after 2–3 weeks, will clinch the diagnosis.

*Scabies*, an infestation with the mite *Sarcoptes scabiei*, commonly produces its characteristic burrows around the wrists and along the sides of the fingers; the diagnosis may be confirmed by direct microscopy of a mite carefully removed with a needle. *Pompholyx* is a form of endogenous eczema presenting with the episodic development of tiny ‘sago-grain’, intensely itchy vesicles on the palms and sides of the fingers. It is commonly triggered by stress or hot weather.

### FITS AND CONVULSIONS

**David Werring**

(See also *TETANY*, p. 671; *TICS*, p. 682; *VERTIGO*, p. 736.)

Epilepsy is derived from the Greek words for ‘taking hold of’. ‘Convulsion’ refers to an intense paroxysm of involuntary muscle contractions, and is not synonymous with ‘fit’ or ‘seizure’, both of which can describe epileptic events involving only alteration of sensation or consciousness. Epilepsy is defined by recurrent epileptic seizures; by definition, a single seizure does not constitute epilepsy. In this section, only epileptic seizures will be discussed, although it should be remembered that other phenomena can mimic epilepsy. In particular, post-syncopal myoclonic jerks following a cardiogenic or vasovagal faint, rigors and tetany can resemble convulsive seizures, so that all episodes of convulsive movements are not necessarily due to epilepsy. Non-epileptic attack disorders (sometimes called ‘pseudoseizures’ or ‘psychogenic seizures’) should also be considered in the differential diagnosis. In describing seizures, ‘tonic’ means sustained muscle contraction, while ‘clonic’ refers to repetitive brief muscle jerks. The cornerstone of the diagnosis of epilepsy remains an accurate witnessed account of the attacks.

Hughlings Jackson, a British neurologist, postulated that epilepsy is due to intermittent dysfunction of the nervous system due to ‘an excessive and disorderly discharge of cerebral nervous tissue [on muscles]’. This fundamental postulate has not been seriously challenged by increasingly sophisticated electrophysiological techniques, which have confirmed the role of hyperexcitable or hypersynchronous cerebral cortex neuronal activity in seizures. Epilepsy is a common chronic condition; indeed, in neurology clinics throughout the UK, one in five referrals is because of epilepsy or suspected epilepsy. Over two-thirds of seizures begin in childhood, with a reduction in incidence after this age, and another increase after the age of 60. In childhood, epilepsy most often has a presently unknown pathological basis and is called ‘primary’, although many cases are likely ultimately to be attributed to a genetic predisposition (e.g. neuronal ion channel dysfunction). Adult-onset seizures are more likely to be due to an identifiable neurological disease (e.g. a brain tumour, a neurodegenerative condition or hippocampal sclerosis). Seizures may be secondary to an intercurrent medical illness (as in sepsis or febrile convulsions, alcohol withdrawal or head trauma).

Seizures may be classified on the above basis (primary or secondary), but can also be classified on the basis of their clinical form (or semiology) that reflects their presumed origin in the brain (generalized – from both hemispheres; or focal – from within one hemisphere) (Box F.11). The major distinction is between partial and generalized seizures. Partial seizures are subdivided into simple (consciousness undisturbed) and complex (consciousness disturbed). Simple partial seizures can be subdivided according to their

<table>
<thead>
<tr>
<th>Classification of epileptic seizures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Generalized (bilaterally symmetrical, no focal symptoms at onset)</td>
</tr>
<tr>
<td>• Tonic, clonic or tonic–clonic</td>
</tr>
<tr>
<td>• Absence</td>
</tr>
<tr>
<td>– Simple – only loss of consciousness</td>
</tr>
<tr>
<td>– Complex – plus brief tonic, clonic or automatic movements</td>
</tr>
<tr>
<td>• Myoclonic seizures</td>
</tr>
<tr>
<td>• Atonic (astatic) seizures with or without myoclonic jerks</td>
</tr>
<tr>
<td>Partial (focal onset)</td>
</tr>
<tr>
<td>• Simple (no loss or alteration of consciousness)</td>
</tr>
<tr>
<td>– Motor – frontal origin</td>
</tr>
<tr>
<td>• Tonic–clonic, Jacksonian, Rolandic, epilepsy partialis continua</td>
</tr>
<tr>
<td>• Sensory (somatosensory, visual, auditory, olfactory, gustatory, vertiginous)</td>
</tr>
<tr>
<td>• Autonomic</td>
</tr>
<tr>
<td>• Psychic</td>
</tr>
<tr>
<td>• Complex (with impaired consciousness)</td>
</tr>
<tr>
<td>• Other</td>
</tr>
<tr>
<td>• Reflex (e.g. photosensitive, reading epilepsies)</td>
</tr>
<tr>
<td>• Febrile seizures of infancy and childhood</td>
</tr>
<tr>
<td>• Hysterical seizures (non-epileptic attack disorder)</td>
</tr>
</tbody>
</table>
FITS AND CONVULSIONS

principal manifestations. Some partial seizures begin with a focal neurological disturbance reflecting the function of the part of the cortex in which the seizure discharge begins (see AURA, p. 41). Partial seizures may also be associated with a transient post-ictal focal disturbance, for example a hemiparesis (Todd's phenomenon). Generalized seizures may be either convulsive or non-convulsive. The most common convulsive generalized seizure is the tonic–clonic (grand mal) seizure. The prototype non-convulsive generalized seizure is the brief loss of consciousness in childhood absence (petit mal). Any partial seizure may generalize to a secondary tonic–clonic seizure.

GENERALIZED SEIZURES

Tonic–clonic seizures (grand mal)

This is the most common form of generalized seizure. There is, by definition, no preliminary focal (partial) aura, although there may be general prodromal symptoms such as fatigue or ill-defined malaise. As the seizure begins, the patient typically cries out during a tonic phase of contraction of extensor muscles of the trunk, limbs and neck (opisthotonus), with subsequent respiratory impairment, laboured breathing noises and cyanosis. There is often reflex emptying of the bladder and occasionally also of the bowel. The patient then enters a clonic phase of generalized symmetrical limb- jerking that lasts for a variable length of time (usually several minutes), followed by deep coma and then a gradual return of consciousness, with post-ictal confusion and disorientation lasting several hours. There may occasionally be automatic behaviour afterwards.

Simple absence (petit mal)

The classic form of childhood generalized seizure is the petit mal seizure. This is a brief absence (usually 10 seconds or less) in which a child loses contact with his or her surroundings. There may be minor myoclonic activity around the eyelids. The attacks start and stop suddenly, they may be very frequent, and the child may not be aware of them. They may present as problems with learning at school. The electroencephalogram (EEG) in classical absence shows characteristic 3-per-second spike-and-wave activity.

Myoclonic jerks

(See CLONUS, p. 101.)
Brief myoclonic jerks occur in numerous epileptic syndromes. They may be associated with an absence, but more commonly they occur without impairment of consciousness. The arms are more frequently involved than the legs.

PARTIAL SEIZURES

Simple partial seizures

Frontal lobe

Frontal lobe seizures are commonly manifest as ‘adversive’ attacks in which there is tonic or clonic deviation of the head and eyes away from the side of the cerebral focus of origin. This is sometimes associated with jerking of the arm or the adoption of a raised flexed posture of the arm on the side to which the head turns. This form of frontal lobe seizure is more common than the classical Jacksonian motor seizure, in which there is a march of movement beginning distally in a digit and spreading up a limb. Both of these forms of motor seizure may be followed by a Todd's paresis, and involvement of the frontal speech areas may cause speech arrest. There may be progression to altered or lost awareness, or secondary generalization into a tonic–clonic generalized seizure. The use of continuous electroencephalographic monitoring during seizures (EEG telemetry) has shown that some frontal lobe seizures cause dramatic motor phenomena including repetitive ‘bicycling’ movements of the legs, sometimes associated with disinhibited behaviour. Unless suspected, these seizure types are easily misclassified as being psychogenic.

Temporal lobe

Seizures beginning in the medial temporal lobe may create sensations of taste or smell (usually unpleasant), epigastric disturbances and pallor, flushing or changes in the heart rate. In addition, there may be psychic phenomena such as déjà-vu and jamais-vu. Seizures originating in the lateral temporal neocortex cause auditory or visual hallucinations, which are often described as having a cinematic playback quality. These focal symptoms often progress to altered or lost awareness (complex partial seizure; see below).

Parietal lobe

These seizures are less common, and may be associated with positive sensory disturbance such as pins and needles, or with distortions of light and colour, often confined to the contralateral half of the visual field. Occipital seizures cause simple, unformed colourful shapes, while more anterior (parietal) sites of origin produce more complex visual phenomena such as patterns or objects.

Complex partial seizures

Complex partial seizures – previously termed ‘psychomotor’ or ‘temporal lobe’ seizures – are differentiated from simple seizures by the impairment of consciousness. Such disturbance of consciousness may be preceded by symptoms of simple partial
type, as above. There may also be contralateral dystonic posturing of the upper limb. Automatic behaviour (automatisms) may occur. Chewing or lip-smacking movements may be seen, as may repetitive fiddling with buttons or clothing. Occasionally, seizures are associated with more complex automatisms, which can lead to problems such as exposure to risk (e.g. climbing out of windows). Very rarely, criminal activity may result, especially in prolonged complex partial status (e.g. shoplifting, vandalism or indecent exposure).

**STATUS EPILEPTICUS**

Most epileptic seizures are self-limiting, but on rare occasions they follow one another in close succession, resulting in convulsive status epilepticus. This is described as a state of recurrent tonic–clonic seizures without recovery of consciousness. The duration most commonly used in defining status epilepticus is 30 minutes, but some experts have suggested a defining period as brief as 5 minutes. It is a medical emergency with a high morbidity and mortality, especially in the elderly. As a first presentation, it is seen in those with symptomatic seizures (e.g. haemorrhagic stroke, electrolyte disturbance or meningo-encephalitis). In patients with a previous diagnosis of epilepsy, status epilepticus is most often seen in severe symptomatic epilepsies and in poorly compliant patients. Most treatment regimens involve early rectal (diazepam) or intravenous benzodiazepines (e.g. lorazepam), followed by intravenous phenytoin. If these standard measures fail, anaesthetic agents and intensive monitoring should be considered at an early stage. It should be noted that non-epileptic attacks constitute a significant proportion of patients admitted to intensive treatment units and, if suspected, require prompt specialist assessment. Absence status is occasionally seen in children, who present with confused behaviour, blinking or small myoclonic jerks. Complex partial status may cause confusion and disorientation, sometimes with automatic behaviour (see above). *Epilepsia partialis continua* is a repetitive rhythmic jerking of a group of muscles (see CLONUS, p. 101). It is most commonly seen in association with frontal lobe cerebrovascular disease.

**INVESTIGATION OF THE EPILEPSIES**

The cornerstone in the diagnosis of epilepsy is a history obtained from the patient and, perhaps more importantly, from witnesses. Epileptic attacks are stereotyped and, on occasion, time is required to obtain accounts of several attacks in order to make an accurate diagnosis. An EEG may add weight to a clinical diagnosis, but can never either prove or disprove epilepsy. The value of interictal recordings is limited, since mild non-specific abnormalities are found in up to 10 per cent of the normal population, while 10–20 per cent of patients with epilepsy do not demonstrate epileptiform abnormalities. The sensitivity of the EEG increases with repeated testing and sleep recordings. Despite its limitations, the EEG remains the most useful single paraclinical diagnostic test for epilepsy. The pattern and location of interictal epileptiform abnormalities not only help to make a diagnosis but also assist in classification of the epileptic syndrome, which is important in determining the prognosis and optimizing treatment. Interpretation of the EEG requires specialized expertise and liaison between the treating physician and neurophysiologist to avoid the over- and underdiagnosis of epilepsy.

A definitive diagnosis of epilepsy can occasionally be made if EEG discharges can be correlated with the patient’s habitual seizures clinically. Continuous long-term EEG recording with video monitoring (EEG telemetry) is expensive and not widely available, but it is the most effective way to evaluate frequent habitual seizures, particularly if a non-epileptic or psychogenic component is suspected. EEG telemetry is used not only for differential diagnosis but also for selecting patients for surgical treatment. A very precise localization of a resectable epileptogenic region is possible with the use of intracranial depth or cortical surface electrodes.

Unless an unequivocal diagnosis of a primary epilepsy is made on clinical and EEG grounds, the use of computed tomography (CT) or magnetic resonance imaging (MRI) is appropriate to look for a treatable structural cerebral lesion, but this will depend on local availability. Imaging, where available, is mandatory in adult-onset patients with focal origin. Once a diagnosis of epilepsy has been made, an attempt should be made to classify the syndrome in terms of seizure type and underlying cause. The number of epilepsy syndromes and causes is formidable, but the fundamental distinctions are between primary and symptomatic epilepsy, and between generalized and partial epilepsy. The primary generalized epilepsies include childhood absence seizures, juvenile myoclonic epilepsy and generalized tonic-clonic seizures in the young adult. Primary partial epilepsies include benign focal motor epilepsy of childhood (*Rolandic epilepsy*) and benign occipital epilepsy of childhood. Symptomatic seizures may be due to a systemic disturbance such as sepsis with fever, hypoxia, hypoglycaemia, electrolyte imbalance or renal, hepatic or respiratory failure. They may be due to toxins, such as drugs (particularly tricyclic antidepressants), alcohol or heavy metals.
FLATULENCE, BLOATING AND ABDOMINAL DISTENSION

Pyridoxine deficiency, porphyria, some inborn errors of metabolism and drug withdrawal are other possible (but rare) causes of symptomatic seizures.

Almost any central nervous system disease may cause symptomatic epilepsy. A brief selection includes: congenital disorders (e.g. tuberose sclerosis, lipid storage diseases, leucodystrophies, Down’s syndrome, microcephaly and hydrocephalus); infective conditions (e.g. meningoencephalitis, especially herpes simplex encephalitis, cerebral abscess, fungal infection and HIV); trauma, diffuse brain injury or penetrating brain injury; cerebral tumours, particularly gliomas, meningiomas and secondary metastases; stroke (infarction, intracerebral or subarachnoid haemorrhage); and degenerative conditions such as Alzheimer’s disease and Pick’s disease. Rarely, multiple sclerosis may present with seizures when a plaque is close to the grey matter of the cortex. Seizures may follow any craniotomy procedure.

The investigations of the various causes of symptomatic seizures may include biochemical and haematological investigation, imaging with CT or MRI, and possibly CSF examination. The clinical situation will dictate which tests are needed.

Non-epileptic attack disorders (which may also be termed pseudo-seizures, psychogenic or hysterical seizures) are increasingly recognized as an important consideration in the differential diagnosis of convulsions. There are some features that raise the possibility of non-epileptic attacks. Even when generalized they will not typically involve self-injury or incontinence. It is usually possible to demonstrate normal pupillary responses, and no focal neurological signs; the heart rate and blood pressure may remain normal, and plantar responses remain flexor. Forced eye closure during the seizure is also a useful pointer towards non-epileptic attacks. Epileptic seizures are stereotyped and usually brief (less than 5 minutes), whereas non-epileptic attacks may change in character and last much longer. An elevation of serum prolactin above 1000 IU on a venous sample taken within 30 minutes of an event (with a normal baseline level) supports the diagnosis of seizure disorder but is not diagnostic. Capture of an event during EEG telemetry is the optimal method of clarifying the diagnosis of non-epileptic attacks, although it must be remembered that epileptic and non-epileptic attacks may co-exist. The management of the patient with non-epileptic seizures is challenging, involving referral to specialist neuropsychiatric services where available. Techniques including cognitive–behavioural therapy may be very helpful, at least in the short term.

FLATULENCE, BLOATING AND ABDOMINAL DISTENSION
Simon Anderson

Wind, abdominal bloating and excessive belching or flatus are common complaints, usually attributed to ‘excess gas’, which is usually incorrect. The volume of gas in the gastrointestinal tract is actually relatively constant (approximately 200 ml) in both fasting and postprandial states and in those with- and without bloating.

Intermittent belching or eructation is commonly caused by certain foods such as onions, cabbage or fatty foods, which delay gastric emptying. Recurrent, prolonged episodes can be caused by the unconscious habit of air-swallowing (aerophagia). It is uncertain if the following other causes can result in bloating as a predominant symptom: infections (giardiasis, Cryptosporidium parvum, Blastocystis hominis and helminths), coeliac disease, disaccharidase deficiency, pancreatic insufficiency, bacterial overgrowth (blind-loop syndrome or strictures) and diverticular disease. An obstruction of the oesophagogastric junction (adenocarcinoma, peptic stricture, achalasia) or gastric outlet (peptic ulcers or tumours) can cause similar symptoms. Delayed gastric emptying without mechanical obstruction, called gastroparesis, is often idiopathic or due to poorly controlled type 1 diabetes. Postprandial gastric bloating due to trapped gas is a well-recognized complication of anti-reflux surgery (fundoplication).

Generalised abdominal daytime bloating is often a feature of irritable bowel syndrome (IBS). Whereas IBS is often associated with altered bowel habit, the Rome III criteria recognize a bloating-predominant form. Recent studies show that this is often due to involuntary incoordinate activity of the diaphragmatic and anterior abdominal wall muscles, rather than an excess of intestinal gas. In cases where bloating is due to excess gas production, this is due to either dietary or bacterial factors. Some foods contain high levels of simple, poorly digested carbohydrates which are fermented by colonic bacteria leading to excess gas production and the formation of osmotically active substances. Reducing these in the diet can have a dramatic effect. Examples of such foods include wheat, onions, broccoli, pulses, beans, stone-fruits, apples, and insoluble fibre. Many patients are helped by avoiding gluten (wheat, barley and rye).

Bacterial factors are poorly understood but thought to be an imbalance in the types of bacteria rather than the overall number. Probiotic medications, used to alter the gut flora, and the use of non-absorbable antibiotics (e.g. rifaximin) are championed, but there is
no conclusive proof of efficacy, and any changes to the gut flora are short term.

Cognitive behavioural therapy has been tried, with good success. The use of smooth muscle relaxants and antidepressants (e.g. low-dose paroxetine) has met with variable success. It is possible that serotonin receptor antagonists/agonists or opiate receptor agonists may be of value.

Premenstrual lower abdominal bloating is a common complaint and ought not to be labelled as IBS. The condition of post-infectious IBS is important to recognize as it can be profound and long-lasting.

Many pathological states can give rise to bloating. When bloating is of recent onset it is important to exclude ovarian cancer and other abdominal malignancies. Other pathological causes include infections (giardiasis, Cryptosporidium parvum, Blastocystis hominis and helminths), coeliac disease, disaccharidase deficiency, pancreatic insufficiency, bacterial overgrowth (blind-loop syndrome or strictures) and diverticular disease. Medications such as lactulose, metformin, bisphosphonates and statins may be implicated in the onset of abdominal bloating.

In the acute setting, a distended abdomen may be due to a mechanical or functional intestinal obstruction. Causes of a mechanical obstruction are numerous and include a volvulus (particularly of the sigmoid colon in the elderly), intussusception (particularly in children), hernia obstruction, tumours, Crohn's disease and adhesions. Functional obstructions include a toxic megacolon (in acute severe colitis) and pseudo-obstruction – including postoperative ileus, hypokalaemia, severe sepsis. Treatment of the underlying cause, for example endoscopic/radiological reduction of a volvulus or correction of an electrolyte imbalance, is of paramount importance. Other measures, such as regularly altering posture (decubitus knee–chin position, upright posture, etc.) or the use of intravenous neostigmine (a parasympathomimetic), may help in toxic megacolon and pseudo-obstruction, respectively.

**FLUSHING**

**Paul Carroll**

Flushing is a slowly spreading erythema of the skin due to a temporary dilatation of the capillaries, conventionally differentiated from emotional blushing only by its severity, duration and extent. The skin of the face, neck and upper anterior chest may be involved. Depending on the underlying condition, flushing may be accompanied by light-headedness, a sense of suffocation, tremors, tinnitus and sometimes nausea and vomiting.

**Blushing** typically results from anger or embarrassment. It produces an instantaneous blotchy or confluent appearance, and symptoms usually date from adolescence.

**Menopausal flushing** (‘hot flushes’) is extremely common in women (approximately 75 per cent of women) at and just after the menopause, but it may occur earlier following bilateral oophorectomy. The flushes, which typically last 1–5 minutes, may be accompanied by sweating and develop spontaneously, sometimes even during sleep. Visible changes occur in 50 per cent of women, and the symptoms typically last 1–5 years, although longer in a minority of women. The mechanism is still unknown but is presumably neurohormonal. Interestingly, one-third of men aged 55–75 report hot flushes; these men often have other symptoms suggestive of a low testosterone concentration, such as decreased muscle strength, lack of energy and low mood. Men receiving hormonal treatment for prostate cancer may suffer from hot flushes.

**Alcohol-induced flushing** can be related to quantity or variety of drink consumed. Large amounts of histamine are found in sherry and some red wines, although none in distilled spirits. Histamine causes flushing, and certain drugs and food may release enough from mast cells to cause a blush. High plasma levels of acetaldehyde, as seen following alcohol consumption in subjects with a deficiency of liver aldehyde dehydrogenase, are likely to contribute to flushing in certain far eastern genotypes. Drugs that cause flushing (particularly in combination with alcohol) are all vasodilators, nicotinic acid derivatives, ciclosporin, sildenafil and similar drugs, chloropropamide, disulfiram, metronidazole and percutaneous absorption of the anti-SCABETIC Tetmosol™ (monosulfiram).

Paroxysmal flushing is the most common clinical feature of carcinoid syndrome and may be associated with migrating wheals. Carcinoid syndrome is due to hepatic involvement by an argentaffin cell tumour, with the production of 5-hydroxytryptamine. Other symptoms are diarrhoea, abdominal pain and right-sided heart failure. In *systemic mastocytosis*, severe flushing attacks, often accompanied by headache, may occur spontaneously or after trauma to skin lesions. Episodes of flushing and diarrhoea may accompany Zollinger–Ellison syndrome, and fainting with flushing can occur with an adrenaline-secreting phaeochromocytoma. It may also occur in insulin-dependent *diabetes* with both hypo- and hyperglycaemia. It may be part of an *epileptic aura.*
Postprandial flushing of the face, especially the nose and adjacent skin, is a characteristic of rosacea. In this disease, reddening later becomes permanent with telangiectasia, as well as papules and pustules (see Fig. E.8). A flushed facial appearance is seen in patients with Cushing’s syndrome, polycythaemia rubra vera and adrenocorticotrophic hormone-secreting bronchogenic tumours (see Fig. S.24).

FOOT, ULCERATION OF

Mark Kinirons

Perforating ulcers of the foot usually occur under the ball of the great toe, but they may affect any pressure area.

Ulcers can form under hard calluses. Anaesthesia appears to be an important precipitating factor, and such lesions are seen in patients with sensory neuropathy of any cause (e.g. diabetes, leprosy, alcoholism and poliomyelitis). Pressure ulcers are also seen in paraplegics. Aggravating factors include foot deformities and ill-fitting footwear that give rise to pressure areas.

Chronic ulceration of the sole can be the presentation of ischaemia (Fig. F.26), and it occurs in those with arteriosclerosis, in heavy smokers and in patients with familial hyperlipidaemia. Vasospasm of the digital arteries occurs in Raynaud’s phenomenon, which can lead to digital ulceration of both the hands and feet. Raynaud’s is characterized by a triphasic reaction of pallor, cyanosis and hyperaemia in response to cold or emotional stress. Cholesterol emboli to the small digital arterioles can present as digital ulceration on the feet, and should be suspected if the patient has recently had angiographical studies/interventions performed. Ulceration of the feet can be the presenting feature of cryoproteinaemia, haemoglobinopathies and hereditary spherocytosis. Intravascular thrombosis possibly plays a role in the development of the ulceration, and the lesions are more common in patients with severe anaemia. In addition, other autoimmune conditions such as systemic lupus erythematosus, antiphospholipid syndrome, scleroderma and other connective tissue disorders should be considered when patients present with ulceration of the lower extremities. Deep skin infections can cause foot ulceration, including fungal (blastomycosis, sporotrichosis and maduromycosis) and bacterial (streptococci, both aerobic and anaerobic; the latter when colonizing an ulcerating expansion of a wound, either traumatic or surgical, is known as Meleney’s ulcer). Mycobacterial infections can present with ulceration of the extremities including the feet (e.g. Mycobacterium marinum found in water and M. ulcerans, responsible for the Buruli/Bairnsdale ulcer, found on grass and introduced into the skin via grass cuts). Rare causes include syphilitic gumma and neoplasms, for example carcinoma cuniculatum, squamous cell carcinoma, malignant melanoma and Kaposi’s sarcoma.

Causes of foot ulceration are listed in Table F.2.

Table F.2 Causes of foot ulceration

<table>
<thead>
<tr>
<th>Ischaemic</th>
<th>Atheroma, diabetes mellitus, Raynaud’s phenomenon, cholesterol emboli</th>
</tr>
</thead>
<tbody>
<tr>
<td>Neuropathic</td>
<td>Diabetes mellitus, leprosy, alcoholism, poliomyelitis</td>
</tr>
<tr>
<td>Vasculitides/ intravascular</td>
<td>Systemic lupus erythematosus, antiphospholipid syndrome, scleroderma, thrombosis cryoproteinaemia, haemoglobinopathies (e.g. sickle-cell disease), hereditary spherocytosis</td>
</tr>
<tr>
<td>Infections</td>
<td>Fungal: blastomycosis, sporotrichosis, maduromycosis</td>
</tr>
<tr>
<td></td>
<td>Bacterial: streptococcal Mycobacterial: M. marinum, M. ulcerans Syphilis</td>
</tr>
<tr>
<td>Malignancies</td>
<td>Carcinoma cuniculatum, squamous cell carcinoma, malignant melanoma, Kaposi’s sarcoma</td>
</tr>
</tbody>
</table>
G

GAIT, ABNORMALITIES OF

Mark Kinirons

Gait may be disturbed by: (i) mechanical defects in the lower limbs and pelvis; (ii) pain in the legs, pelvis and lower lumbar spine; (iii) disease of the muscles; (iv) disease of the nervous system – increased tone, either basal ganglial or pyramidal; weakness, either pyramidal or peripheral; ataxia, either sensory or cerebellar; and cortical; (v) disease of the vestibular apparatus; and (vi) hysteria.

MECHANICAL

Inequality in the length of the legs, congenital dislocation of the hips, ankylosis of the knee or hip joints, and deformities of the feet give rise to a characteristic bold, painless limp, the source of which is readily found on examination of the limbs.

Coxa vary and coxa valga may lead to characteristic gaits. Painless ankylosis of both hips leads to all movements being made at the knees, ankles and feet, giving a short-stepping smooth gait, almost as if the patient were on roller-skates. This is seen, for instance, in some cases of ankylosing spondylitis.

Painful limp, due to pain in the pelvis or lower limb, is easily recognized by the manner in which the patient puts the weight on the sound leg and hurries off the affected one. The source of the pain may be in the limb itself, or it may be referred from disease in the pelvis, lumbar spine or cauda equina. Localized pain usually means localized disease at that site, referred pain tending to have a more diffuse and linear distribution, but many exceptions occur. The pain referred from a diseased hip may be felt only in the knee – an important feature in children with tuberculosis of the hip – and occasionally a pain root is limited to a small area in the foot (SI) or the lateral border of the leg (L5).

Local tenderness at the site of the pain often indicates local disease, but it may equally well be present in referred pain. On the other hand, local deformity or swelling always means disease at that point. The more important causes of a painful limp are the following:

- The joints:
  - Injuries
  - Arthritis or arthrosis of lumbar spine, or of the hip, knee, ankle and/or foot on one or both sides
- The bones:
  - Injuries
  - Neoplastic, congenital or metabolic disease
  - Inflammatory infective or degenerative disease

DISEASES OF THE MUSCLES

Diseases of muscles, although rare, can cause characteristic disturbances of gait. In the heredofamilial myopathies, usually seen in early life, the gait is waddling, the muscles are weak and either hypertrophied or atrophied, and sensation is normal. In muscular dystrophy (MD), the muscles weaken, leading to an abnormal gait. The severity depends on the type of MD, Duchenne’s MD being the worst and leading to death prematurely in early adult life. In myotonia congenita, members of affected families experience from birth a peculiar difficulty in relaxing the muscles after voluntary contraction. Thus, on attempting to walk, the muscles go into a tonic spasm, but this can be worked off by continued exercise. Diagnosis is made on the family history, the presence of prolonged contraction after voluntary effort, the production of a persistent localized contraction on percussion of the affected muscles, and high-frequency discharges in the electromyogram, likened on the loudspeaker to the noise of a dive-bomber. A similar myotonia is seen in dystrophia myotonica, a familial disease usually of adult life but occurring also in children; the gait is disturbed by myotonia, but weakness and atrophy of the quadriceps and of the dorsiflexors of the feet are a further embarrassment to walking. The presence of wasting and myotonia in the face, sternomastoids and forearm, and the frequent presence of premature baldness, cataracts and testicular atrophy, indicate the correct diagnosis. In myasthenia gravis, the legs, in common with the rest of the musculature, may fatigue rapidly; the gait is normal after rest, but as fatigue supervenes, it becomes shuffling, unsteady and weak.

The weakness and extreme hypotonia of the muscles in amyotonia congenita interfere with gait in those children who survive the first critical years of infancy. They learn to walk later, and they then present the unsteadiness of weakness, but even this incapacity may be outgrown.

DISEASE OF THE NERVOUS SYSTEM

Spasticity due to bilateral pyramidal disease will affect both legs, as in congenital spastic diplegia,
spinal cord compression, multiple sclerosis, subacute combined degeneration of the cord in the early stages, intramedullary tumours and syringomyelia. Tone is increased in the extensors and adductors, so that the limb is held in extension, with plantar flexion of the foot, and some degree of adduction. The gait is stiff, the toes scrape the ground and, if the adduction and spasticity are severe, there is a ‘scissors gait’. Weakness increases the disability. A unilateral pyramidal lesion gives rise to a similar stiff, extended limb, which is dragged around its normal fellow by tilting the pelvis, thus overcoming the adduction and allowing the flexed foot to clear the ground.

The rigidity of extrapyramidal disease affects the extensors and flexors equally, but the legs are held slightly flexed at the hip and knee because the flexors are more powerful than the extensors. Steps are short and shuffling. The patient tends to walk faster and faster, as if chasing his centre of gravity. If pushed backwards, he tends to run backwards with short hasty steps. Fixity of expression, flexion of the neck and trunk, adduction of the arm with flexion of the elbows and the characteristic rhythmic tremor of the forearms, hands and fingers afford diagnostic assistance when the extrapyramidal lesions are due to Parkinson’s disease, one of the earliest signs being a failure to swing the arm on walking. A similar gait results from extrapyramidal lesions due to parkinsonism. Parkinsonism is seen with drug-induced, arteriopathic cerebral degeneration, carbon monoxide poisoning, hepatolenticular degeneration and encephalitis lethargica, and (rarely) after severe head injuries, such as from boxing.

Weakness plays a part in pyramidal lesions, but it is often difficult to distinguish the relative importance of this weakness and the associated spasticity in the disturbances of gait that are described above. On the other hand, weakness due to the anterior horn cells or of the peripheral nerves to the legs gives rise to abnormal gaits, the features of which depend on the distribution of the weak or paralysed muscles. Where there is foot drop, as in any form of polyneuritis, injuries to the common peroneal nerve, poliomyelitis or a lesion of the cauda equina, there is a high-stepping gait, in which the foot is lifted high and then slapped down on the ground. If the calf muscles are paralysed, as in a lesion of the posterior tibial nerve, the gait loses its natural spring. Furthermore, disease affecting the motor fibres often attacks the sensory fibres too; proprioceptive sensory loss then adds a sensory ataxic element to the gait, which becomes clumsy, unsteady, irregular and broad-based as in tabes, many types of polyneuritis and gross disease of the cauda equina. When sensory ataxia thus complicates muscular weakness, balance is worse in the dark or when the eyes are shut.

Sensory ataxia has been mentioned as a factor in the clumsy, incoordinated, noisy, wide-based gait of polyneuritis, tabes, lesions of the cauda equina and some cases of subacute combined degeneration of the cord. A second form of ataxic gait is seen as a result of disease or injury of the cerebellum or its connections. The gait is wide-based and clumsy, but it is little aggravated by darkness or by closing the eyes. There is a tendency to deviate towards the side of the lesion, but overcompensation may occur, with consequent deviation to both sides in an irregular, staggering and drunken manner. The normal ‘swing’ of the arm on the affected side may be diminished or lost, but this feature is often absent. ‘Cerebellar’ ataxia is seen in multiple sclerosis, the heredo-familial ataxias, occasional cases of tabes without proprioceptive sensory loss, and inflammatory, neoplastic, degenerative, traumatic and vascular lesions of the cerebellum.

Disease that affects the cortex can frequently cause abnormal gait. ‘Marche à petits pas’ is used to describe a gait pattern due to multiple cortical infarcts. A form of disconnection seems to impair higher cortical connections. Advanced dementia frequently impairs gait, but the diagnosis is obvious by then.

VESTIBULAR DISEASE
Disease of the labyrinth, the vestibular nerve or the vestibular nucleus in the pons can give rise to disturbances of gait. Vertigo makes the patient feel disoriented; the gait is unsteady, and there is a tendency to deviate to the affected side. This occurs in acute phases of Ménière’s disease, acute labyrinthitis and vascular lesions of the pons.

HYSTERIA
Hysteria is sometimes responsible for abnormal gaits. There is no set pattern, but, however bizarre hysterical gaits may be, they have in common a certain improbability and flamboyance, and a tendency to subside gracefully and safely on the floor, and to stagger in the direction of objects upon which to lean. ‘Astasia abasia’ is a term that was formerly used for inability to stand or walk despite normal movements of the limbs when recumbent, but it is desirable to recall that, in hereditary spinocerebellar ataxia, and in affections of the flocculonodular lobe and the vermis of the cerebellum, a gross ataxia of gait may be found despite good performance in all tests of coordination when in bed. This is due to the fact that these functions,
of balance and coordination, are subserved by different regions of the cerebellum, which can be involved separately in disease processes, coordination depending on the integrity of the lateral lobes of the cerebellum and balance on the midline structures. Hysterical gaits are recognized by their inconsistencies, and by a quality that can best be described as insincerity; confirmation is to be found in the presence of psychoneurosis and the absence of organic disease. Unlike organic disorders of gait, they can sometimes be cured by suggestion.

GALLBLADDER, PALPABLE
Harold Ellis

PHYSICAL SIGNS
On occasion, a grossly distended gallbladder in a thin subject may be visible as a distinct globular swelling in the right upper abdomen. However, palpation is the physical method of examination in detecting enlargement of the gallbladder. One may feel an oval, smooth swelling moving downward close behind the anterior abdominal wall when the patient inspires, descending either from beneath the right costal margin near the tip of the ninth rib, or attached to the undersurface of a palpable liver in the right mid-clavicular line. As it enlarges, the tumour generally extends inwards as well as downwards so that it may ultimately cross the midline below the level of the umbilicus. It may be large enough to be palpable bimanually in a thin patient, but it does not fill out the loin in a way that a renal tumour may do. It may or may not be tender, depending on whether the cause of the enlargement is or is not associated with inflammation. It feels firm and tense rather than hard. An impaired but not quite dull note is obtained on percussion.

DIAGNOSIS FROM OTHER SWELLINGS
Palpable gallbladder must be distinguished particularly from four groups of conditions:

- From carcinoma arising in the bile ducts or gallbladder itself
- From tumours in or attached to the liver in the neighbourhood of the gallbladder – secondary neoplasm (Fig. G.1), primary hepatoma or more rarely gumma, abscess or hydatid cyst
- From mobile kidney, hydronephrosis or renal tumour
- From tumours in the neighbouring organs, such as carcinoma of the pyloric antrum or the right suprarenal gland

Clinical features, as described below, will often enable an accurate diagnosis to be made. These may be supplemented by appropriate radiological studies and by ultrasound or by retrograde percutaneous transhepatic cholangiography (PTC) (Fig. G.2) and, if necessary, with fine-needle aspiration.

Carcinoma of the gallbladder
It is often difficult to decide whether a tumour is merely an enlarged gallbladder or a growth infiltrating and replacing it, since in either case there may be a history extending over years of gallstones, with biliary colic,
pyrexia and even jaundice, and primary new growth of the gallbladder is usually associated with gallstones. The rapidity of the enlargement in the absence of any definite cause will suggest growth, particularly in a person of an age with a higher risk of cancer. Careful palpation may show that the mass is not smooth, as in the case of most simple gallbladder enlargements, but more or less nodulated or covered with bosses or irregularities, which in themselves suggest new growth. In some cases, there may be secondary deposits in the liver and ascites, and sometimes the enlargement of the left supraclavicular lymph nodes points to malignant disease with metastasis. Notwithstanding these points, the differential diagnosis may be so difficult that further special investigations will be necessary for decision.

**Tumours attached to or in the liver**

Those most likely to be mistaken for enlargement of the gallbladder are Riedel’s lobe, secondary carcinoma of the liver and, much more rarely, hepatoma, gumma, abscess or hydatid cyst. It may, by physical examination, be impossible to distinguish a Riedel’s lobe from an enlarged gallbladder or from a mobile kidney. Speaking generally, a Riedel’s lobe (a congenital abnormality of no other clinical significance) usually descends from the liver farther to the right than does a gallbladder, and it is more apt to simulate an enlarged or a mobile kidney.

**Metastatic deposits** in the liver nearly always cause considerable (and sometimes enormous) enlargement and great hardness of the organ, not infrequently associated with jaundice. The diagnosis depends first upon the discovery of a primary growth, which in the case of carcinoma is likely to be in the large bowel, stomach, pancreas or breast, or, in the case of melanoma, the eye; and second, on the discovery in the liver of several separate nodules, some of which may be felt to be umbilicated, that is to say depressed in their central part and raised around the edges.

**Hepatoma**, although rare in the UK, occasionally occurs in patients with cirrhosis and may be multifocal. In the Far East and in eastern Africa it is far more common, and, in patients from those areas, is an important condition to consider in differential diagnosis.

**Gumma** of the liver is rarely encountered nowadays, and when it occurs is usually mistaken for new growth, unless there is a convincing history of syphilis or the effects of tertiary lesions are visible elsewhere – especially gummatous lesions of the skin or leucoplakia of the tongue. The diagnosis may be confirmed by obtaining a positive serological reaction, or by the beneficial effects of antisyphilitic treatment, although this does not always lead to rapid disappearance of a gumma of the liver. Even when the liver is inspected at laparotomy, the diagnosis between gumma and new growth is not always easy.

**Abscess** of the liver, if it is to simulate an enlargement of the gallbladder, is likely to be a single large abscess that, if it has not arisen in some pre-existent mass (such as a gumma, new growth or hydatid cyst), is almost certain to have been acquired in a tropical country where the patient has suffered from amoebic dysentery. The diagnosis may not be evident until the mass is punctured with an exploring needle, yielding the typical ‘anchovy sauce’ pus.

**Hydatid cyst** of the liver is seldom situated in such a position as to cause difficulty of diagnosis for gallbladder enlargement. More usually, the cyst is embedded in the liver substance or projects from its upper surface. The diagnosis might be entertained if the patient is known to have hydatid cysts elsewhere or came from an area where this disease is endemic. However, in most cases, it is suggested by ultrasonography or computed tomography of the liver, and it is sometimes determined only when laparotomy has been performed. It might have been suggested by the discovery of eosinophilia, and also by the specific hydatid serum reaction if the hydatid cyst is alive and active. But latent or calcified hydatid cysts cause no symptoms, do not produce an eosinophilia and are not associated with a positive hydatid blood–serum reaction. Their walls, if calcified, can be seen on radiographs of the region.

**Distinction between an enlarged gallbladder and a mobile kidney or hydronephrosis**

There may be no jaundice to suggest gallbladder trouble, nor need there be any urinary changes to suggest kidney, so the diagnosis may have to be made chiefly by palpation. Facts to stress are that a gallbladder is more easily felt anteriorly than posteriorly, while the reverse is the case with the kidney; that the kidney is, as a rule, the more freely movable of the two; that it is seldom possible to demarcate the upper pole of an enlarged gallbladder in the way that the top of a movable kidney can sometimes be defined; that with kidney tumour the loin is dull, while with gallbladder enlargement it is resonant; and that, on rather firm bimanual palpation, the patient may experience a peculiar sickening sensation that is characteristic of kidney. In cases of doubt, ultrasound or an intravenous pyelogram will demonstrate whether or not the right kidney is normal (see also KIDNEY, PALPABLE, p. 334).
Tumours of other organs simulating enlargement of the gallbladder
These may be distinguished to some extent by the fact that new growths of the pylorus, transverse colon or suprarenal that are big enough to simulate an enlargement of the gallbladder seldom have the smooth oval outline that the gallbladder nearly always possesses. In addition, there may have been symptoms attributable to the primary growth, such as dilatation of the stomach, coffee-grounds vomit, or evidence of secondary deposits in the liver, in the left supraclavicular lymph nodes or elsewhere to indicate the diagnosis.

Modern imaging techniques (ultrasound and computerized axial tomography, CAT) can usually provide anatomical delineation of an enlarged gallbladder and differentiate the other masses enumerated above. Nevertheless, in some of these cases, it is impossible to exclude enlargement of the gallbladder without resorting to laparotomy.

CAUSES OF GALLBLADDER ENLARGEMENT
These include the following:
- Empyema of the gallbladder
- Chronic pancreatitis
- Carcinoma of the head of the pancreas
- Cholecystitis as a result of:
  - Gallstones
  - New growth
- Typhoid fever
- Obstruction of the common bile duct by a gallstone
- Obstructed of the cystic duct by a gallstone
- Simple mucocele

It is noteworthy that gallstones comparatively seldom lead to enlargement of the gallbladder. If the associated inflammation does not progress to empyema, the gallbladder usually becomes thick-walled, contracted and embedded in dense adhesions that prevent it from dilating, even when the cystic or common bile ducts become obstructed by a stone. Indeed, in a middle-aged patient in whom there has not been any very definite attack of biliary colic, the occurrence of progressive and considerable enlargement of the gallbladder, associated with deepening jaundice and without ascites, arouses serious suspicion of a lesion of the head of the pancreas that has extended along the pancreatic duct so as gradually to occlude the common bile duct (Fig. G.3). The most common cause of these symptoms is either chronic pancreatitis or carcinoma of the head of the pancreas or of the ampulla of Vater. In obstruction of the common bile duct due to gallstones, the gallbladder is as a rule not palpable; in obstruction due to carcinoma of the head of the pancreas, it is usually distended and is palpable in about 50 per cent of patients (Courvoisier’s law, which states that ‘In the presence of jaundice, a palpably enlarged gallbladder is unlikely to be due to stone’) (Fig. G.4). Painless progressive jaundice suggests a carcinoma arising at the ampulla of Vater and, if this ulcerates, the stools may be positive for occult blood. Jaundice preceded by epigastric or upper lumbar pain is more likely to be due to carcinoma or chronic pancreatitis of the body of the pancreas. Sometimes, sloughing of part of the tumour allows the pent-up bile to escape into the duodenum, with puzzling temporary remission or even disappearance of the jaundice.

In cases where gallstones are the cause of the enlargement, there is nearly always tenderness over the gallbladder and pain when it is palpated firmly, associated with a rise in temperature, possibly with rigors, especially if the inflammation has

Figure G.3 Computed tomography scans showing a cancer in the head of the pancreas (white arrow) causing obstruction of the bile ducts (red arrow) and gallbladder (blue arrow).
spread to the bile ducts (infective or suppurative cholangitis). Leucocytosis, with a relative increase in the polymorphonuclear cells, would indicate that, in addition to gallstones, there is empyema of the gallbladder demanding urgent surgical treatment. Another cause of empyema of the gallbladder, albeit rare, is typhoid fever. The diagnosis is not difficult as a rule, for in most of the cases there will be no question of new growth or of gallstones and the patient will have been suffering from a prolonged asthenic fever that has already been diagnosed serologically. In some typhoid patients, bacillary infection of the gallbladder causes it to enlarge rapidly, even to the extent of rupturing spontaneously and causing general peritonitis. In less severe cases, the inflammatory products discharge themselves naturally by the bile passages.

Simple mucocele of the gallbladder is a relatively unusual event that results from the impaction of a gallstone at the outlet of the gallbladder when it happens to be empty. The walls of this organ continue to secrete mucus so that it becomes greatly distended with perfectly colourless mucoid liquid, free from bile pigment. The fluid is sterile. There are usually no symptoms. Such a mucocele may be mistaken for a mobile kidney. Usually, the differential diagnosis can be established by radiological examination (cholecystography or intravenous urography) or by ultrasound or computed tomography. However, the diagnosis of the nature of the mass is sometimes obscure until revealed by operation.

**GANGRENE, PERIPHERAL**

Harold Ellis

Gangrene results from death of a part of the body or an organ as a result of it being deprived of its blood supply (Fig. G.5), and with superadded bacterial infection of the resultant dead tissues. Ischaemia without infection results in a sterile infarction. In this section, we consider gangrene of the limbs, which is seen much more frequently in the toes and foot than in the fingers and hand, although the same pathological conditions apply, in general, to both.
The list of possible aetiologies is extensive, but in the great majority of patients encountered in the Western world with this condition, the cause is arteriosclerotic disease, which may or may not be complicated by diabetes mellitus.

CAUSES OF PERIPHERAL GANGRENE

These are listed in Box G.1.

Trauma
The diagnosis of traumatic gangrene can rarely give rise to any difficulty, in that the clinical features of history and examination will betray the cause. Inadvertent intra-arterial injection of a barbiturate or other irritant drug is a rare anaesthetic accident, but is seen more frequently in drug addicts; the additional factor in the latter cases is that the injected material is often also contaminated with bacteria.

Frostbite
Frostbite affects subjects exposed to intense cold without adequate protection, and results from ice crystals forming in the tissues, followed by capillary sludging and thrombosis in the small vessels.

The affected skin of the fingers or toes becomes first cold, white, immobile and anaesthetic. As the tissues rewarm, the affected skin becomes red and painful, blisters develop and minor trauma allows ingress of bacteria, with resultant gangrene of the dead tissues.

Infection
Gas gangrene results from Clostridium perfringens (welchii) and other Clostridium species. The organism is a Gram-positive, spore-forming bacillus that is an obligatory anaerobe and produces powerful exotoxins. The organisms are found in soil and faeces. Typically, gas gangrene is an infection of deep, penetrating wounds where there is extensive devascularized muscle, which acts as a perfect culture medium for the organism. It is particularly seen in war wounds, but sometimes involvement of the abdominal wall or cavity may follow operations upon the alimentary tract. Gas gangrene may complicate the amputation of an ischaemic limb.

The incubation period is about 24 hours. The patient becomes severely toxæmic, with rapid pulse, shock and vomiting. The temperature is first elevated and then becomes subnormal. The affected tissues are swollen, and crepitus is palpable due to liberated gas from protein destruction. The overlying skin becomes gangrenous, and infection spreads along the muscle planes, producing first dark red, swollen muscle and then frank gangrene of the infected muscle from origin to insertion.

Synergistic gangrene
This condition, also known as progressive bacterial gangrene, is caused by the synergistic action of two or more organisms, commonly aerobic haemolytic Staphylococcus and microaerophilic non-haemolytic Streptococcus. When this occurs around
the perineum or buttocks, or following abdominal surgery, coliform bacteria may also be present. It is more common in diabetics and immunosuppressed patients, and is often related to recent trauma or surgery (where it was previously termed ‘progressive postoperative gangrene’) or infection (e.g. an ischio-anal abscess). Where it affects the scrotum, it has been termed ‘Fournier’s gangrene’.

Around the wound or infection, an area of cellulitis appears, which spreads rapidly. The area is exquisitely tender, and the subcutaneous tissues slough with an offensive odour and most often the presence of gas (which can be mistaken for gas gangrene). The overlying skin becomes gangrenous, and the patient becomes profoundly septic.

There is a high mortality rate, and the condition can only be treated by a combination of high-dose, broad-spectrum antibiotics together with radical excision of the affected area (which may often have to be repeated on several occasions).

**ARTERIAL DISEASE**

**Arteriosclerosis**

This is by far the most common cause of peripheral gangrene of the toes and foot (but occasionally the fingers) in the Western world. The patient is more often male than female, usually over the age of 50, and invariably a long-standing and heavy cigarette smoker. There is a history of progressively deteriorating claudication leading to rest pain, with cramping calf pain experienced even at rest in bed. There is then some incident of minor trauma that may require direct questioning to elicit, such as nicking the toe while cutting the nail, or a minor blow to the foot, or pressure from a tight shoe, which enabled organisms to invade the ischaemic tissues through an often negligible wound. A gangrenous patch on the heel or over the lateral malleolus may result from pressure on these areas from a hard mattress.

Clinical examination of the leg reveals, apart from the obvious area of gangrene, that the peripheral skin is cold and pale or cyanosed. Elevation of the leg increases the pallor, while depression of the leg over the side of the bed usually produces a deep cyanosis (Buerger’s test). The peripheral pulses are absent.

A general assessment of the patient is, of course, mandatory and often reveals features of general ischaemic disease, such as a history of angina or of previous coronary ischaemic episodes, a previous stroke or transient ischaemic attacks.

It is important to investigate the presence of co-existing diabetes mellitus. This will exacerbate the seriousness of the condition (see Diabetic gangrene, below).

The important radiological investigation is arteriography, which defines the extent of the disease and also enables the surgeon to decide whether vascular reconstruction is possible by surgery or balloon angioplasty (Fig. G.6). Duplex sonography is now replacing arteriography in many centres. This technique takes longer to perform and is more subjective, but it is non-invasive and can provide better information on the significance of stenoses.

**Thrombo-angiitis obliterans (Buerger’s disease)**

This condition, which some regard as a variant of arteriosclerotic disease of the limbs (‘juvenile arteriosclerosis’), but which the majority of experts consider a special entity, is almost confined to men in their 20s and 30s who are very heavy cigarette smokers. It was originally thought to occur only in Jews of eastern European extraction, but it is now known to be widely distributed and is seen, for example, quite commonly in Chinese and Arab individuals, and men from the Indian subcontinent.

The distal arteries of the lower (and often the upper) limbs are affected, and show round-cell infiltration and intimal proliferation with intraluminal thrombosis. Adjacent veins and nerves become involved in this inflammatory process, and there may

![Figure G.6 Volume-rendered computed tomography angiogram of the aorta (green arrow) and distal vessels showing heavily calcified superficial femoral arteries bilaterally (white arrowheads) and acute occlusion from thrombus on the left (blue arrow).](image)
be associated superficial venous thrombosis. The gangrenous process, which usually affects both feet (and often the fingers), is preceded by a history of claudication, eventually with rest pain, and sometimes by episodes of superficial or deep vein thrombosis in the legs. Unlike arteriosclerosis, the femoral (and often also the popliteal) pulses are usually present, although the dorsalis pedis and posterior tibial pulses are lost. Late cases of this condition are pitiful; progressive gangrene may have led to serial amputations of all four limbs.

Arteriography shows a fairly typical appearance of relatively normal arteries down to the proximal part of the brachial or popliteal, with distal vessel obliteration. Biopsy of an occluded artery or vein may provide histological confirmation of the disease.

**Embolus**

Gangrene due to embolism will be sudden in its inception, and rapid in its onset. Potential sources of emboli include (Fig. G.7):

- Left atrium – atrial fibrillation with mitral stenosis, atrial myxoma
- Heart valves – endocarditis affecting diseased valves
- Left ventricular wall – mural thrombus after myocardial infarction or from ventricular aneurysm
- Aorta – from aneurysm or arteriosclerosis
- Interventricular septum – paradoxical embolus via a septal defect originating in the systemic veins (rare)

Among this list, the most common causes of arterial emboli are dislodgement of a mural clot from a myocardial infarct, often occurring around 10 days previously, and from clot in the left atrium in atrial fibrillation.

The history is usually one of sudden pain in the limb, which soon becomes white and cold. Sensation may disappear, and the muscles become rapidly paralysed. Over the next few hours, the limb becomes anaesthetic, and skin staining appears that does not blanch on pressure. If untreated, peripheral gangrene will ensue. Pulses below the block, which usually takes place at a major vessel bifurcation, including that of the aorta, are absent.

Paradoxical emboli are uncommon. In patients with a persistent foramen ovale or other septal defects, a clot originating in the deep veins of the leg or pelvis may not only impact in the pulmonary arterial tree but also pass across the septal defect and lodge in the arterial system. This is particularly likely to occur after a pulmonary embolus, as the raised pulmonary artery pressure results in increased shunting across a septal defect if this is present.

A limb embolism must be differentiated from acute thrombosis occluding an arteriosclerotic artery, where there is more likely to be a history of preceding claudication. Both may require urgent surgery, but an embolus is treated by balloon embolectomy, whereas acute thrombosis in arteriosclerosis may necessitate urgent endarterectomy or a bypass procedure. An arteriogram is necessary to differentiate between the two, the filling defect in an embolism showing a smooth rounded outline like a cigar butt, whereas that of acute thrombosis is irregular and merging indefinitely with the jagged outline of the locally diseased vessels.

**Diabetic gangrene**

Diabetes mellitus, if severe and poorly controlled, may result in infective gangrene of the foot (Fig. G.8). This results from a combination of diabetic microangiopathy, with subintimal arteriolar thickening producing vascular impairment, an increased susceptibility to tissue infection as a result of reduced host defences, and an associated peripheral neuropathy (often of ‘glove and stocking’ distribution), which renders the soft tissues more susceptible to trauma. Osteomyelitis of the bones of the foot is a common complication of this combination of infection and neuropathy. Since this is a disease of small vessels, the peripheral pulses are palpable. Those at the ankle may be difficult to detect because of oedema of the tissues, but their patency can be confirmed by means of a Doppler probe.
In young patients, this may well be the only pathology producing the gangrene. In older subjects, however, with co-existent arteriosclerotic disease, the presence of diabetes adds to these additional factors listed above to produce a much poorer prognosis than in patients with otherwise uncomplicated arteriosclerotic occlusive disease in the limb.

Raynaud’s disease and Raynaud’s phenomenon
Raynaud’s disease is caused by spasm of the small arteries in the digits in the hands (and often the feet) of unknown aetiology, which was described by Maurice Raynaud, a physician in Paris, in 1862. It is almost confined to young females in their teens and twenties. As a result of exposure to cold, the fingers become white, and later slate-blue (see Fig. F.19, p. 202). As the hands warm, they change from livid purple to deep red, and this cycle can be precipitated by plunging the hands first into a basin full of cold and then into one of hot water. Many attacks do not pass through all these colour changes, with either pallor or cyanosis being the predominant feature. Because the disease affects only the terminal vessels, the radial pulse is normal and, in those cases where the toes are affected, the ankle pulses are not lost. As might be expected, the disease is subject to exacerbations in the winter and remissions in the summer, but with gradual progression. With more advanced cases, chronic paronychia, with ulceration and gangrene of the fingertips may be seen, although this is much less common than in Raynaud’s phenomenon (see below). Raynaud’s phenomenon – sometimes termed ‘Raynaud’s syndrome’ or ‘secondary Raynaud’s disease’ – is the term applied to a similar clinical picture affecting the fingers, but in which there is an underlying organic disease of the arteries of the upper limb (Fig. G.9). The list of such diseases is extensive and includes:

- Trauma
  - Penetrating or closed arterial injury
  - Frostbite
  - Persistent use of vibrating tools
  - Arterial thrombosis following inadvertent intra-arterial injection of barbiturate, etc.
- Arterial disease
  - Arteriosclerosis
  - Thrombo-angiitis obliterans (Buerger’s disease)
  - Cervical rib with embolism
- Connective tissue disease
  - Scleroderma (systemic sclerosis)
  - Systemic lupus erythematosus
  - Rheumatoid arthritis
- Drugs
  - Beta-blockers
  - Ergotism

Among this list, patients with scleroderma are particularly likely to progress to digital gangrene (see also FINGERS, DEAD (WHITE, COLD), p. 201).

Ergotism
Long ago, epidemics of painful digital gangrene were seen called ‘Saint Anthony’s Fire’. This is now known to be due to the ingestion over a long period of rye bread contaminated with Claviceps purpurea, which produces ergot, which causes small arterial spasms and eventual intimal proliferation. Today, ergotism is seen occasionally in patients with severe migraine who take excessive amounts of ergotamine tartrate. The author has seen only one example of this, in an elderly lady.
Venous gangrene
Gangrene is an unusual complication of thrombosis of the deep veins of the leg. This has to be extremely extensive, involving the femoral, iliac and pelvic veins. The legs are grossly swollen, oedematous and cyanosed. Gangrene may be limited to the toes but may spread to the feet and even to involve an extensive area of the limb (Fig. G.10).

Grip, Disturbances of
Mark Kinirons
Man’s privileged position in the animal kingdom is attributable to his prehensile upper limbs, as well as to his superior brain. Our capacity to grip enables us to lift, carry, climb and handle tools. In terms of functional anatomy, grip depends primarily on the long flexor muscles of the fingers, and the opposition of the thumb to the other digits. The latter movement is possible because the metacarpal bone of the thumb lies in a plane at right angles to that of the other metacarpals, and because the carpometacarpal joint of the thumb has such a wide range of movement. Flexion thus carries the thumb medially across the palm, in opposition to the other fingers.

Grip is impaired by any lesion affecting the motor supply of muscles of the forearm and hand. Thus, upper motor neurone lesions in corticospinal pathways, radicular lesions affecting the C7, C8 and T1 roots, and peripheral nerve lesions may all result in weakness of grip (see MONOPLEGIA, UPPER LIMB, p. 417). As regards single peripheral nerve lesions, it is noteworthy that although median or ulnar nerve palsies can affect the grip, the greatest disability results from an isolated radial nerve palsy. This is because without fixation of the wrist in extension, the grip becomes ineffective.

Apart from purely motor lesions, grip can also obviously be impaired by severe incoordination, dystonia, involuntary movement or sensory deprivation. It may also be affected by disease of the joints or muscles, although, in general, muscle disease tends to be of proximal rather than distal distribution. Myotonia dystrophia is characterized by a normal grip, but a delayed relaxation of the grip.

Groin, Swelling in
Harold Ellis
Swellings in the groin are a common clinical problem, and examples are likely to be encountered in every general surgical outpatient clinic, as well as being found on routine examination of patients in other departments. The great majority can be diagnosed accurately by careful history and clinical examination.

Anatomy
The groin is not a well-defined anatomical zone but may be divided into the inguinal and femoral regions by the inguinal ligament (not the groin skin crease) (Fig. G.11). The inguinal ligament runs from the easily defined anterior superior iliac spine to the pubic tubercle at the lateral extremity of the pubic crest. The pubic tubercle can be felt with relative ease in the thin subject; in those who are more obese, it can be defined as the bony prominence that can be felt at the apex of the tendon of adductor longus with the hip in the flexed and abducted position. Swellings above the medial part of the inguinal ligament can be described as inguinal, and those below as femoral.

Figure G.10 Gangrene due to extensive venous thrombosis may involve an extensive area of the limb.

Figure G.11 Anatomy of the groin.
CLASSIFICATION

As there are numerous potential causes of a swelling in the groin, a two-stage mental process is required in making a differential diagnosis: first, what anatomical structure is involved; and, second, what pathological entities may arise therefrom (Box G.2).

By far the most common are: inguinal hernia in male adults and children; inguinal and femoral hernia in adult females; inguinal lymphadenopathies; and saphena varix.

SKIN AND SUBCUTANEOUS TISSUES

Sebaceous cysts occur in the groin but are less common than on the scrotal skin.

Lipomas represent by far the most common swellings arising from the subcutaneous tissue, and they may be found in the inguinal canal (lipoma of the cord), although these latter usually merely represent herniating extraperitoneal fat. Lipomas show the classical features of a soft, lobulated, mobile, fluctuant and transilluminable mass. The much rarer liposarcoma is rapidly growing, vascular and invasive.

Neurofibroma may occur unusually as a solitary swelling and is suggested by accompanying café-au-lait pigmented areas of skin. More often, multiple lesions are found here and elsewhere as part of von Recklinghausen’s disease.

LYMPH NODES

(See also LYMPHADENOPATHY, p. 394.)

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<thead>
<tr>
<th>Box G.2</th>
<th>Classification of groin swellings according to tissues of origin</th>
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<tr>
<td><strong>The skin and subcutaneous tissues</strong></td>
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<tr>
<td>• Sebaceous cyst</td>
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<td>• Lipoma</td>
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<td>• Neurofibroma</td>
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<td><strong>Lymph nodes</strong></td>
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<td>• Lymphadenopathy due to infection, secondary neoplasm, lymphoma</td>
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<td><strong>Blood vessels</strong></td>
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<td>• Ectopic testis</td>
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<td>• Lipoma of the cord</td>
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<td>• Hydrocele of the canal of Nuck in the female</td>
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<td><strong>Psoas sheath</strong></td>
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<td>• Psoas abscess</td>
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Lymphadenopathy

The groin lymph nodes drain the lower limb, the external genitalia (scrotum and penis in the male, labia and vulva in the female), the lower anal canal, the buttock and the abdominal wall below the level of the umbilicus. All of these areas must be searched carefully for a primary focus of sepsis or neoplasm in a patient with groin lymphadenopathy.

Groin lymphadenopathy is usually multiple, but a large single node in the region of the femoral canal (Cloquet’s node) may mimic an irreducible or strangulated femoral hernia. The common causes of groin lymphadenopathy are listed in Box G.3.

Inflammatory lymphadenopathy

There is usually a history of an inflammatory lesion involving the tissue drained by the lymph nodes. A search is made for a septic focus on the foot or toenails, between the toes, and on the leg, buttock, external genitalia, lower abdomen and anal canal. This may appear trivial, and the lymphadenopathy may continue for up to 3–4 weeks following resolution of the initial cause. In the early phase, the node may feel indurated and tender, after which it becomes firm and softens as it resolves. Reactive lymphadenopathy may be single or multiple, but the nodes usually feel discrete. Should resolution fail to occur over 3–4 weeks, biopsy becomes important to identify specific infections or malignancy.

The inguinal lymph nodes enlarge bilaterally with a penile or vulval syphilitic chancre. They remain discrete and do not suppurate. This is in contrast to the groin lymphadenopathy in chancroid, where the enlarged groin nodes have a marked tendency to ulcerate (see PENILE LESIONS, p. 502).

A range of viral infections will cause generalized lymphadenopathy, but when sustained over weeks with ill-health and malaise, the Paul–Bunnell antibody reaction may be requested where infectious mononucleosis is suspected. In modern practice, toxoplasmosis and other causes of lymphadenopathy related to AIDS should be considered. In these cases the

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<th>Box G.3</th>
<th>Causes of groin lymphadenopathy</th>
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<td><strong>Inflammatory</strong></td>
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<td>• Acute – from a septic focus in an area drained by groin nodes</td>
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<td>• Part of generalized inflammatory lymphadenopathy (viral) [e.g. glandular fever, AIDS]</td>
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<td><strong>Neoplastic</strong></td>
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<td>• Metastatic from squamous carcinoma or melanoma in an area drained by groin nodes</td>
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appearance and feel of the lymph nodes are similar to that with other reactive lymphadenopathies, but in AIDS lymphadenopathy is more common in the neck and axillae.

**Metastatic carcinoma or melanoma**

Firm, hard nodes that grow into a confluent fixed mass as the disease progresses are typical of metastatic carcinoma, which may be confirmed by either biopsy or needle aspiration cytology. If the primary has not been identified, a careful search must be performed for primary lesions in the territory drained by the relevant lymphatics. This includes a rectal examination with proctoscopy and sigmoidoscopy, careful examination of the perineum and scrotum, and a thorough search of the legs, the feet and between the toes.

**Lymphoma**

The clinical features of both Hodgkin's and non-Hodgkin's lymphomas are similar, but in the latter condition the nodes may be hard and more closely resemble those of metastatic carcinoma. In Hodgkin's lymphoma, the nodes often have a typical rubbery feel and may be matted together such that a clinical diagnosis can be quite confident. The diagnosis is confirmed by biopsy, which should include an intact node taken (if possible) from a region other than the groin. Once the diagnosis has been confirmed, the extent of the disease should be fully identified (staging) by further investigations, which include chest X-ray and computed tomography scanning of the chest and abdomen.

**VASCULAR SWELLINGS**

Vascular swellings in the groin should be easily recognized, yet the saphena varix is often mistaken for a femoral hernia. Aneurysms of the common femoral artery are relatively rare, and are most frequently seen following previous arterial surgery as a pseudoaneurysm (Fig. G.12). Occasionally, an inflamed pseudoaneurysm or mycotic aneurysm may present in a drug addict who has used the femoral artery for vascular access. The diagnostic importance of this is that it may be mistaken for an abscess, with catastrophic results when drainage is attempted.

**Saphena varix**

**Anatomy**

The saphena varix develops when there is incompetence of the sapheno-femoral junction and saphenous system. The proximal part of the saphenous system becomes dilated where reflux of turbulent blood impinges on the anterior wall of the vein. It lies in the upper femoral triangle just below the usual location for a femoral hernia, and just medial to the femoral pulse.

![Figure G.12 Pseudoaneurysm of the left femoral artery following an aorto-bifemoral bypass using a Dacron® graft. The earlier left below-knee amputation was performed for extensive gangrene of the foot.](image)
Anatomy
The common femoral artery lies immediately below the mid-point between the anterior superior iliac spine and the mid-point of the pubic symphysis. Aneurysms rarely extend into the profunda or superficial femoral artery.

Clinical features
In the absence of inflammation, the diagnosis is obvious, with easily palpated expansile pulsation. A thrill may be palpable, and auscultation often demonstrates a bruit, indicating turbulent blood flow within the aneurysmal sac. A rapidly expanding pseudoaneurysm, or more particularly mycotic aneurysm, may be mistaken for an abscess. Gentle palpation through the induration will always reveal substantial and abnormal pulsation. Occasionally, there may be the swelling of the distal limb due to compression of the vein or lymphatics.

THE HERNIAL ORIFICES

Body wall
Groin hernias are the most frequent type of hernia and may give rise to a number of acute complications. They are distinguished by simple clinical features readily identified on examination.

Inguinal hernia
The inguinal hernia is by far the most common variety of hernia. Approximately 3–5 per cent of the population have some variety of hernia, and 70–75 per cent of these are inguinal. Inguinal hernias are very much more common in men, and they represent over 90 per cent of all hernias in men. In women, inguinal hernias are also nearly twice as common as femoral hernias, although there is a misconception that, as femoral hernias commonly occur in women, they are also more common than inguinal hernias.

Most groin hernias present as a soft lump in the groin that may be associated with local pain or discomfort on standing or lifting, or after heavy work. The clinical features of an inguinal hernia are of a lump arising just superior to the lower one-third of the inguinal ligament. Most hernias are reducible, in which case there will be an impulse on coughing. In small hernias, the sensation of reduction is perhaps the most reliable clinical sign confirming the presence of a hernia.

Anatomy
The inguinal canal is formed by the descent of the testis from the posterior abdominal wall during early development. During fetal development, it passes obliquely through the anterior abdominal wall, extending from the internal inguinal ring at the mid-point of the inguinal ligament to the external inguinal ring just above the pubic tubercle. The anterior wall is formed by the external oblique aponeurosis and incorporates the lower-most fibres of the internal oblique at the internal ring. Inferiorly, there is the inguinal ligament, and posteriorly the transversalis fascia, reinforced medially by the fibres of the conjoint tendon. The internal oblique and transversus muscles arch over the canal superiorly. Indirect inguinal herniae arise through the internal ring, passing into what is probably a congenital sac consisting of a persistent patent processus vaginalis. Direct inguinal herniae occur as a result of weakness in the posterior wall of the inguinal canal medial to the inferior epigastric artery, which passes in the medial border of the internal ring.

Clinical features
The vast majority of patients notice a lump in the groin, which may be asymptomatic or cause an aching or dragging sensation (Fig. G.13). Occasionally, the presenting symptoms are those of small-bowel obstruction when the bowel herniates and then becomes stuck in the sac, with oedema and obstruction at the hernia ring. A local, irreducible and acutely inflamed lump implies strangulation of the hernial contents, which may be extraperitoneal fat, omentum or intestine.

Inguinal herniae in infants and children
These are almost invariably indirect, with 90 per cent occurring in boys. The lump may be noticed by the mother while bathing the child, or when the child is straining, coughing or upset. It may also be detected either during the postnatal examination or as the child first starts to walk. It is frequently difficult to demonstrate the hernia in the clinic, but a clear history from the mother is sufficient to make a confident diagnosis. As there is a definite risk of strangulation, the surgeon should not refrain from

Figure G.13 Right inguinal hernia.
exploring the affected side where an appropriate and reliable history can be obtained. The differential diagnosis consists of a hydrocele of the cord, undescended testis or torsion of the testis. Careful examination of the scrotum is therefore mandatory.

**Inguinal hernia in adults**
Essentially, adults may have either indirect or direct inguinal hernias, with sliding hernias representing a variety (usually indirect) in which an adjacent partly peritonealized organ herniates as part of the hernia sac wall. This usually involves the caecum, the sigmoid colon or the bladder.

**INDIRECT INGUINAL HERNIAS**
These arise from the internal ring lateral to the inferior epigastric vessels and pass down the inguinal canal within the cord. Frequently, they pass down into the scrotum and may reach a substantial size (Fig. G.14). This variety of hernia occurs ten times more frequently in men than in women and is more common than the direct variety. It is identified by an impulse passing obliquely from lateral to medial as the hernia is produced on coughing, and by the sensation of a reduction in the line of the inguinal canal on gentle pressure. Once the hernia is reduced, it can be controlled by digital pressure over the internal ring, even with the patient standing or coughing.

**DIRECT INGUINAL HERNIA**
This variety of hernia is extremely rare under the age of 40 years and typically presents in the elderly. A bulge is identified over the most medial part of the inguinal ligament that may reach sizeable proportions but very rarely descends into the scrotum. It is uncommon for direct hernias to obstruct, and they usually have an obvious and prominent cough impulse that appears to come directly out through the body wall rather than obliquely in the line of the inguinal canal.

The differential diagnosis of inguinal herniae includes hydrocele of the cord, herniated extraperitoneal fat or lipoma of the cord, inguinal lymphadenopathy, ectopic testis and femoral hernia. The inguinal hernia is best distinguished from the femoral hernia by carefully identifying the pubic tubercle, where the insertion of the inguinal ligament can usually be felt. The inguinal hernia arises above the inguinal ligament, passing over this point towards the medial side. The femoral hernia, which may appear to fold over the inguinal ligament, can be felt to arise in its deeper part from the femoral canal below and lateral to a finger on the pubic tubercle.

**Femoral hernia**
Although relatively common as a cause of small-bowel obstruction, femoral hernias are less common than inguinal, incisional and paraumbilical hernias, representing only 3-5 per cent of all hernias. They are two to three times more common in women than men, with femoral hernias representing more than 20 per cent of all hernias in women. They occur more frequently on the right side, and over 30 per cent present with a complication such as obstruction or strangulation.

**Anatomy**
The femoral hernia descends through the femoral canal bordered medially by the lacunar ligament, posteriorly by the ilipectineal band, and anteriorly by the inguinal ligament. As these three rigid walls limit femoral canal expansion, which can only occur by displacing the femoral vein laterally, there is a definite risk that the hernia will become irreducible or strangulate. The sac then descends to the sapheno-femoral junction, where the least resistance turns the hernia forwards and upwards to lie anterior to and sometimes even extended above the inguinal ligament (Fig. G.15).

**Clinical features**
Femoral herniation is rare before late adult life. It may present as a firm or even rubbery lump in the groin that often represents no more than herniated extraperitoneal fat, together with the tissue accumulated around the sac as it descends through the femoral canal and up through the cribriform fascia. As a result,
the hernial sac can almost invariably be palpated in the upper medial femoral triangle, even when the contents are reduced. It is unusual to feel a cough impulse in a femoral hernia unless it contains omentum or gut that is freely reducible. An irreducible hernia, particularly in the presence of strangulation, will be tender, tense and often inflamed. On careful palpation of the pubic tubercle, the neck of the sac may be felt below and lateral as it passes under the inguinal ligament.

THE TESTICULAR APPARATUS
An ectopic testis may be palpated in the region of the external inguinal ring, superficial to the inguinal ligament, or rarely at the root of the penis. It is the size and consistency of the testis, and gives a testicular-type sensation on compression. The important clue to the diagnosis, of course, is that the testis on the side of the swelling is absent from the scrotum.

A hydrocele of the cord presents as a cystic swelling at the external inguinal ring. Traction on the testis draws the swelling downwards with the cord – a diagnostic physical sign.

A lipoma of the cord as it emerges from the inguinal canal does not demonstrate a cough impulse. However, it frequently co-exists with an indirect inguinal hernia sac, representing merely a collection of extraperitoneal fatty tissue on the sac wall.

In the female, a hydrocele of the canal of Nück (a cyst of the processus vaginalis along the round ligament) is a common differential diagnosis of an irreducible inguinal hernia. It is smooth, painless, fixed in position, fluctuant, and transilluminates brilliantly.

THE PSOAS SHEATH
Pus within the psoas sheath, originating from advanced tuberculous disease of the lumbar spine, from fistulating Crohn's disease or, rarely, from an appendix abscess, may track beneath the inguinal ligament and present in the groin.

GUMS, BLEEDING
Leandros Vassiliou & Mark McGurk

DEVELOPMENTAL LESIONS
The oral cavity may be involved by haemangiomas or vascular lesions and, on occasion, lymphangiomas. All of these lesions are liable to bleed with trauma, and they are particularly hazardous following dental extractions, should these involve the bone of the mandible or the maxilla. They are not normally a diagnostic problem.

DENTAL DISEASES
Bleeding from the gums is a common complaint, the predominant underlying local cause being periodontal disease, which is almost endemic within the human race. The common causative factor is poor oral hygiene. Bacterial biofilms adhere to tooth surfaces becoming organized into dental plaque and calculus. Bacterial toxins induce an inflammatory response causing gingivitis rendering the mucosa oedematous and hyperaemic. Any minor trauma to this swollen and inflamed mucosa, such as mastication or tooth-brushing, causes bleeding. Pregnancy with the associated hormonal changes may exacerbate a pre-existing gingivitis and pregnancy granulomas can occur (Fig. G.16).

If left untreated, the inflammatory process becomes destructive with the result that periodontal disease ensues. The combination of localized bacterial infection and an aggressive immune response by the host causes destruction of the periodontal ligament and alveolar bone supporting the teeth; pockets then form between the gum and the tooth root, where further bacteria accumulate. There may also be a purulent discharge from...
the necks of the teeth, and halitosis. Slow progressive loss of the alveolar bone causes gingival recession, and progressively loosening and loss of teeth. Over the age of 20 years, periodontal disease is the commonest cause of tooth loss rather than dental decay.

Dental caries, when present, may be obvious to the naked eye, or less conspicuously between the teeth (interdental caries) or beneath the gingival mucosa (root caries). This also allows an accumulation of bacteria, leading to localized infection and generating a vicious circle of gingival inflammation, hyperaemia and haemorrhage. If both of these conditions are left untreated, abscess formation will eventually occur, which is discussed elsewhere (see JAW, SWELLING OF, p. 314).

**INFECTION**

The oral cavity is commonly involved in bacterial infection due to dental causes. However, generalized infection of the oral mucous membrane due to bacteria is rare, the majority being caused by viruses.

**Bacterial infection**

The most dramatic example of bacterial infection is acute necrotizing ulcerative gingivitis (ANUG), which is characterized by a proliferation of Vincent’s organisms, *Treponema vincentii* and *Fusiformis fusiformis*. It is not a common condition and is usually associated with a terrible standard of oral hygiene or a person who is immune-suppressed. It has been linked to stressful life events as occur in war zones or national disasters that lead to social deprivation. The predominant features are bleeding of the gums, soreness and halitosis. There is a characteristic ulceration, necrosis and blunting of the interdental papillae with considerable debris and slough in the gingival crevice. There may be an associated fever, malaise and regional lymphadenitis. It may also be a manifestation of reduced resistance to infection as in acute leukaemia, agranulocytosis or sequelae of cytotoxic chemotherapy.

In the African continent, the immune status of the patient may be drained by measles and the chronic effects of malnutrition, and this can lead to a disease called cancrum oris. Small areas of dark necrosis occur in the cheeks or lips, which rapidly progress to produce widespread necrosis and destruction of the circumoral structures (Fig. G.17).

**Viral infection**

Acute inflammation of the oral cavity occurs with viral infections, the most common being herpes simplex and herpes varicella-zoster. The primary infection with herpes simplex may often be subclinical; if not, acute herpetic gingivostomatitis is encountered. This is characterized by severe inflammation and vesicle formation, leading to ulceration of the entire oral mucous membrane with constitutional symptoms of fever, malaise and enlargement of the regional lymph nodes. The gingivae are acutely inflamed, swollen and bleed easily. The disease is usually self-limiting after a few days, and only symptomatic relief and the maintenance of oral hygiene are required. Secondary infection of the vesicles may be prevented by chlorhexidine or tetracycline mouthwashes.

Reactivation of the herpes zoster virus (shingles) in the distribution of the trigeminal nerve may cause severe erythema, confluent vesicle formation and ulceration in the exact distribution of the involved branch of the trigeminal nerve (usually the maxillary branch). This is not a diagnostic problem as mucosal lesion invariably occur with skin vesicles. The condition is again self-limiting, and the intraoral lesions require only symptomatic relief, although an antiviral agent may be valuable in the protracted case.

The Epstein–Barr virus (infectious mononucleosis) may cause a gingivostomatitis in which the gingivae are characteristically red, swollen and haemorrhagic, similar to the appearance found in acute leukaemia or scurvy. Acute bacterial ulcerative gingivitis may follow.
A high proportion of patients show petechiae in the soft palate, which, because of its characteristic appearance, is of diagnostic importance.

Patients with HIV infection may develop significant inflammation of the gingival tissues not commensurate with the level of oral hygiene and periodontal disease. They are also prone to acute necrotizing ulcerative gingivitis and acute necrotizing periodontitis.

**BLOOD DYSCRASIAS**

Abnormal or defective bone marrow activity, as in aplastic anaemia or leukaemia, can present with swollen and bleeding gums due to abnormalities of granulocytes, and low platelets.

In acute leukaemia, the swelling is also due to the local accumulation of leukaemic cells in the gingival tissues. Thrombocytopenia can give rise to intra-oral purpura in the palate as it is subject to trauma during mastication. In the buccal and labial mucosa, where the surface epithelium is not so tightly bound down to the underlying connective tissue, large blood blisters may form.

Haemophilia and von Willebrand’s disease are associated with troublesome bleeding after dental extraction unless appropriate measures are taken.

**DISORDERS OF BLOOD VESSELS**

Scurvy can be conveniently discussed under this heading, as the main defect is one of abnormal collagen production leading to capillary fragility. There is also a general lack of tissue resistance, which, in the gingival mucosa, may lead to superimposed infection.

It is probably this latter aspect that produces most of the gingival enlargement characteristic of the disease, as in the presence of good oral hygiene the swelling is far less marked. Although scurvy is now an uncommon disease in the developed world, it can still be found among old and neglected people with a restricted diet, and also in people who for dietary reasons reduce their input of food containing ascorbic acid. It may also be found in alcoholics and patients with peptic ulceration existing on a milk diet.

Hereditary haemorrhagic telangiectasia (also known as Osler-Weber-Rendu syndrome) (Fig. G.18) is caused by a capillary abnormality, and the head and neck region is a common site. In the mouth these abnormal vessels are visible through the mucous membrane, and minor trauma may produce a persistent haemorrhage. This may be difficult to control in hidden areas such as the nose. Von Willebrand’s disease and Henoch–Schönlein purpura may also present with oral manifestations, but these are not a predominant feature of the disease.

**MALABSORPTION**

Patients with malabsorption may show signs of anaemia in the mouth, with a red and swollen tongue and pallor of the mucosa. In addition to this, the gingival mucosa may haemorrhage readily.

**CHEMICAL POISONING**

Mercury was previously used in the treatment of syphilis coining the expression ‘a night with Venus and a life time with mercury’. The drug produces a severe acute stomatitis with profuse salivation, halitosis.
and painful swellings of the lips, gums, tongue and cheeks. The patient may become manic (mercury was used in the hat trade, hence the term ‘mad as a hatter’), and may develop a metallic taste and cervical lymphadenopathy. This method of treatment has long been discontinued, but poisoning can still occasionally occur from industrial exposure. Teeth loosen and gums become hyperaemic and tend to bleed, which, as always, is aggravated by poor oral hygiene. Eventually, necrosis of the gingival mucosa may occur.

Phosphorus was historically associated with ‘phossy jaw’ (similar to bisphosphonate osteonecrosis), not infrequently ending in death as the result of fatty degeneration of the liver and heart. Since restrictions are now laid upon the use of crude yellow phosphorus in the manufacture of matches, this condition is now almost unknown. Consequently, modern exposure comes with suicide attempts using rat or other vermin poisons that contain phosphorus.

Arsenic and lead are rare causes of gingival bleeding and usually arise from industrial contamination. The gingivae are again inflamed and swollen and bleed easily; in the case of lead or bismuth, there is a characteristic blue line at the gingival margin known as the Burtonian line. Cisplatin, a chemotherapeutic drug can induce a similar picture. Other signs of poisoning may be present, particularly pigmentation of the skin, vomiting, diarrhoea, and hyperkeratosis of the soles of the feet and the palms of the hands. Generalized peripheral neuritis may be found in the case of arsenic, and the symptoms given under anaemia in the case of lead. Arsenic may be found in excess in the hair, or lead may be detected in the faeces or in the urine.

IATROGENIC

Overdose with warfarin may produce spontaneous bleeding of the oral mucosa, and this will require vitamin K preparations or transfusion of fresh-frozen plasma to control the haemorrhage. The toxic effect of cytotoxic agents on the bone marrow may produce a clinical picture similar to those seen with blood dyscrasias.

GUMS, HYPERTROPHY OF

Leandros Vassiliou & Mark McGurk

True hypertrophy of the gingival mucosa is relatively rare, and accordingly enlargement can conveniently be classified into that caused by a cellular infiltration and that caused by predominantly fibrous tissue. It should be remembered that the most common cause of gingival enlargement is infection associated with the dental structures (for discussions, see GUMS, BLEEDING, p. 226; JAW, SWELLING OF, p. 314).

EPULIS

The term ‘epulis’ denotes a swelling arising from the gum, most frequently either a pyogenic granuloma or fibroepithelial polyp or rarely a peripheral giant-cell granuloma. The swellings may be sessile or pedunculated in shape and covered by pink or red mucosa. Histologically, they exhibit a core of granulation or fibrous tissue covered by epithelium.

The pyogenic granuloma, fibrous epulis (Fig. G.20) and fibroepithelial polyp represent an exaggerated soft-tissue response to minor trauma, and are treated by simple excision and curettage. Denture granulomas are histologically similar to fibroepithelial polyps, but arise from low-grade persistent denture trauma. Pseudofolds of mucous membrane with a fibrous tissue stroma are formed in the region of a traumatic denture flange (Fig. G.21). Treatment is again by excision, with attention to the prosthesis.

Infiltration of the gingivae by angiomatous tissue occurs as part of a haemangioma (see JAW, SWELLING OF, p. 314), and should be distinguished from a localized proliferation of capillaries due to chronic irritation, such as that which is caused by a carious tooth or a retained dental root. The latter will resolve with removal of the irritant stimulus, while the former is usually part of a more extensive proliferation involving adjacent structures.

PREGNANCY

Generalized swelling of the gingival mucosa may occur, but the classical pregnancy granuloma is usually confined to one interdental papilla (Fig. G.22). Histologically, it is an exaggerated reparative response and the tissue consists of immature granulation tissue. Management is driven by the symptoms. Normally the lump is unsightly and it may bleed intermittently so most patients request surgical excision, which has
the advantage of a specimen for histological analysis. Good oral hygiene measures contain the generalized condition, which then recedes after pregnancy.

GIANT-CELL GRANULOMA
The aetiology of this lesion is again unknown. It arises in the young age group, and is confined to the tooth-bearing areas of the jaws, with the mandible as the most frequent location. The peripheral giant-cell granuloma (Fig. G.23) arises from the gingival margin as a localized fleshy mass, which bleeds easily. The treatment is surgical excision. Histologically, the picture is of vascular fibrous tissue impregnated with multinucleated giant cells.

BLOOD DYSCRASIAS
Acute leukaemia and scurvy were discussed previously (see GUMS, BLEEDING, p. 226). In acute leukaemia gingival masses may arise as a result of leukemic cells migrating into the tissues, forming leukemic deposits. Gingival masses can also be an extranodal manifestation of lymphomas and normally pose a diagnostic problem. They grow quickly and are destructive in nature (Fig. G.24).

Wegener’s granulomatosis is a disease of focal necrotizing vasculitis that affects the upper and lower respiratory tracts. Occasionally, a proliferative gingivitis may occur that arises interdentally and spreads mainly along the buccal gingivae. Extensive periodontal destruction may occur, and the diagnosis can only be obtained by biopsy. Pointers are associated chest symptoms and renal disease with a significantly raised ESR.

FIBROUS HYPERPLASIA
Hereditary gingival fibromatosis (Fig. G.25) is a rare autosomal dominant condition in which all the gingival tissues become enlarged to such an extent that the

Figure G.21 Denture granulomas (a) that have occurred due to continual irritation from a loose-fitting denture (b).

Figure G.22 Pregnancy tumour.

Figure G.23 Peripheral giant-cell granuloma.
teeth may become almost buried, and in the child will interfere with tooth eruption. Mucosal inflammation is not always a feature, and treatment is by surgical reduction and the maintenance of good oral hygiene. There may, in addition, be hirsutism and thickening of the facial features, associated epilepsy and mental retardation.

IATROGENIC FIBROUS HYPERPLASIA

In a mouth with poor hygiene and pre-existing inflammation, certain drugs such as phenytoin (used to control epilepsy), cyclosporine and calcium-channel blockers (predominantly nifedipine) may cause gingival swelling. It differs from gingivitis with inflamed bleeding gums, but it is characterized by pale, firm enlarged gingivae that may obscure the crowns of the teeth (Fig. G.26). Treatment is through scaling and polishing of the teeth and the maintenance of good oral hygiene, but in some cases it may be necessary to withdraw the drug or excise the hyperplastic tissue.

GUMS, RETRACTION OF

Leandros Vassiliou & Mark McGurk

PERIODONTAL DISEASE

Retraction of the gums is commonly associated with chronic periodontal disease. The accumulation of dental plaque results in inflammation and destruction of the alveolar bone that supports. Normally the gum retracts as the bone recedes. So common is this process with age that it is often referred to as ‘getting long in the tooth’. Vigorous attention to oral hygiene will limit the rate of bone loss, but in some immunologically susceptible patients the condition is progressive, with little evidence of infection. Gingival retraction may be a manifestation of HIV infection.

MALIGNANCY

Gingival margin leukoplakia (gingival margin keratosis) (Fig. G.27) is a rare condition that should be considered a premalignant condition and requires high-level vigilance by the clinician. The leukoplakia migrates along the attached gingiva and replaces the periodontal ligament causing tooth loosening. After loss of the teeth there is delayed healing of the socket. The skill is to ascertain the point where the process has accelerated and become invasive.

Other malignant conditions such as sarcomas (Fig. G.28) can also cause retraction of gums as a result of their biological nature and destructive potential.

EOSINOPHILIC GRANULOMA

The solitary eosinophilic granuloma produces an area of bone loss in the jaw, and there is an associated soft-tissue swelling with gingival ulceration. It is now accepted as a member of the myeloma family. The condition may present with loosening of the teeth, the failure of a tooth extraction site to heal, or a pathological fracture. The jaws may also be affected by multifocal eosinophilic granuloma where, with loosening of the
teeth, there is also generalized inflammation of the oral cavity and gingival enlargement. Management is still evolving and advice on treatment should be sought from a medical oncologist.

RADIOTHERAPY

Radiation to the oral cavity in the management of malignant tumours induces a severe mucositis of the oral mucosa, which is characterized by erythema, superficial ulceration and pain. Irradiated jaw bone, particularly if accompanied by chemotherapy, has reduced resistance to infection as well as limited regenerative capacity. If the mucosal barrier is breached, the underlying bone becomes infected and the mucosa retracts away from the necrotic bone. Such is the evolution of osteoradionecrosis (Fig. G.29).

The condition is all but intractable. Early lesions may respond to prolonged courses of pentoxiphylline and Vitamin E (tocopherol), but the established lesion is resistant to simple measures. Eventually pain and fistulae drive the patient to surgery which is fraught with complications. The object is prevention with good oral hygiene, and careful dental care, both prior to and after treatment.

GYNAECOMASTIA

Paul Carroll

Gynaecomastia is benign enlargement of the male breast due to an increase in the duct tissue and in the periductal stroma. It is a common clinical occurrence seen in up to two-thirds of pubertal males and up to 30 per cent of adults. It is thought to arise as a result of an imbalance between the stimulatory effects of oestrogen and the inhibitory effect of free androgen on the breast tissue. If the condition lasts for longer than a year, the stroma becomes fibrous and the gynaecomastia tends to persist. The condition should not be confused with fat in the mammary region; in this case, no glandular tissue is palpable behind the areola. Other conditions that cause swelling in the breast should be excluded, such as carcinoma, lipoma or neurofibroma. Mammography and breast ultrasound are useful in investigation.

Gynaecomastia is common in neonates and at puberty. When it occurs in older age groups, a pathological cause is more likely, although in the elderly it may be a physiological accompaniment of declining testicular function.

The causes of gynaecomastia are detailed in Box G.4.

NEONATAL GYNAECOMASTIA

Approximately 70 per cent of male neonates have some breast enlargement. In just over half of these cases, fluid (‘witch’s milk’) can be expressed
on squeezing. The breast enlargement is probably due to the effect of placental oestrogens and human chorionic gonadotrophin (hCG) stimulating the Leydig cells of the baby’s testes to produce oestrogen.

The witch’s milk may possibly be the result of maternal prolactin. Histological examination shows the typical features of a lactating breast.

**PUBERTAL GYNAECOMASTIA**

The vast majority of boys develop a minor degree of gynaecomastia at the time of puberty (Fig. G.30). Before the gynaecomastia appears, there is a rise in plasma oestradiol that anticipates the expected pubertal rise in plasma testosterone. Along with the increased oestradiol, there is an increase in prolactin, but the levels fall as the gynaecomastia develops. In boys in whom pubertal gynaecomastia does not occur, this sequence of hormonal events does not take place. In some boys, the gynaecomastia is more marked, and may even approximate to the normal female breast. Often, the condition arises in one breast only, or develops on one side some weeks or months before it appears in the other. Occasionally, fluid can be expressed. In an appreciable number of cases, there is a history of neonatal gynaecomastia, or even a family history, suggesting that there may be a constitutional sensitivity to oestrogen secreted by the Leydig cells of the testis. This sensitivity could possibly be mediated by increased numbers of oestrogen receptors in breast tissue.

Mild, early gynaecomastia usually regresses, but moderate to severe degrees tend to persist. Although, in theory, anti-oestrogens such as tamoxifen should be
of value, they seem to have little effect on anything but mild gynaecomastia. The patient is best referred early for plastic surgery. A periareolar incision should leave a virtually invisible scar.

It is important to consider other causes of gynaecomastia in a pubertal boy, such as drug ingestion, and to examine the testicles carefully for the presence of a tumour. If one testis appears to be normal and the other one is small, the ‘normal’ testis may be harbouring a tumour. The oestrogens produced by the tumour may have suppressed pituitary gonadotrophins and caused failure of development of the contralateral testis. Testicular ultrasonography is a useful way of detecting small tumours.

SENESCENT GYNAECOMASTIA

Occasionally, patients in the sixth or later decades of life may develop gynaecomastia (Figs G.31 and G.32). There may be associated loss of libido and occasional hot flushes. The plasma testosterone level is reduced, while plasma oestradiol levels remain normal, although plasma oestrone levels rise due to increased conversion from androstenedione. Sex hormone-binding globulin (SHBG) levels rise, with a consequent increase in the binding of testosterone. The result is a further reduction in free testosterone and an imbalance in the ratio of free testosterone to free oestrogens. The serum gonadotrophin levels are raised.

Pathological gynaecomastia is found in men where there is a disturbance in the normal balance between testosterone and oestrogen function. This results from conditions where there may be a decrease in testosterone production, a decrease in testosterone action, an increase in oestrogen production, or an increase in oestrogen activity. It should be noted that gynaecomastia is not usually a feature of hypogonadism secondary to pituitary or hypothalamic disease.

CONDITIONS ASSOCIATED WITH DECREASED TESTOSTERONE PRODUCTION

Congenital

Klinefelter’s syndrome describes a phenotypical male with 47,XXY sex chromosome genotype. The condition presents in adolescence, with seminiferous tubule dysgenesis giving rise to primary hypogonadism and male infertility. There is clinical evidence of gynaecomastia in a significant number of cases, along with small atrophic testes and aspermatogenesis. There is a decrease in testicular production of testosterone, with a consequent increase in the oestrogen:androgen ratio.

Congenital anorchia describes phenotypically normal 46,XY males with failure to locate the testes on
surgical exploration. Around 50 per cent of these patients develop gynaecomastia due to a low production of testosterone.

**Defects in testosterone synthesis** due to enzyme deficiencies in the metabolic pathway from cholesterol to testosterone result in incomplete virilization of the male fetus during embryogenesis. ‘Male pseudohermaphroditism’ is a term used to describe individuals who have testes and an XY chromosomal constitution but ambiguous external genitalia. Examples of enzyme deficiencies include congenital adrenal hyperplasia (20, 22-desmolase deficiency, 3-beta-hydroxysteroid dehydrogenase deficiency or 17-alpha-hydroxylase deficiency), failure of conversion of testosterone to dihydrotestosterone because of 5-alphareductase deficiency, or failure of testosterone production because of 17-beta-hydroxy-steroid dehydrogenase deficiency. The clinical manifestations vary widely from hypospadias to grossly abnormal appearances with a small penis exhibiting chordee, a bifid scrotum, a persistent urogenital sinus, and a vagina opening into the posterior urethra. A rudimentary uterus and Fallopian tubes may be present. The testes are undescended or present in the labioscrotal folds. Gynaecomastia may appear at puberty.

**Acquired**

*Bilateral testicular atrophy* is associated with decreased production of testosterone. This occurs in the presence of normal production of oestrogens (oestriadiol and oestrone) from extraglandular sources, resulting in a reduction in the androgen: oestrogen ratio. Gynaecomastia is a common feature.

The most common causes of testicular atrophy acquired after puberty are the following: (i) viral orchitis due to mumps virus, echovirus or group B arbovirus; (ii) trauma; (iii) neurological disease (e.g. myotonic dystrophy, traumatic paraplegia, syringomyelia or Friedreich's ataxia); (iv) granulomatous disease involving the testis (e.g. leprosy); and (v) renal failure – gynaecomastia is present in approximately 50 per cent of patients receiving haemodialysis.

**Conditions associated with resistance to testosterone action**

**Congenital**

*Androgen resistance* describes either testicular feminization syndrome or Reifenstein's syndrome. In these conditions, the underlying pathology is a resistance at the androgen receptor to endogenous and exogenous androgens. The result is a range of incomplete virilization in a 46,XY sex chromosome genotype, resulting in:

- Phenotypic women with testicular feminization (male pseudohermaphrodite) associated with complete resistance at the androgen receptor. The appearance is female, there is absent pubic and axillary hair, the external genitalia resemble those of a normal female, but there is a blind vagina. The testes are usually intra-abdominal but may lie in the inguinal canal or in the labia majora. The breasts are those of a normal female, except that the nipples and areolae are often small.
- Complete resistance at the androgen receptor, examples of which include phenotypic men with hypospadias and gynaecomastia (Reifenstein's syndrome) and infertile male syndrome.

**Conditions associated with increased oestrogen production**

**Testicular tumours**

Testicular tumours such as those of the stromal cells (e.g. Leydig cells and Sertoli cells) produce gynaecomastia because of increased oestrogen secretion. The other germ cell tumours (e.g. chorionic carcinoma, embryonal carcinomas and teratomas) do so because human chorionic gonadotrophin (hCG), produced by the tumour, stimulates testicular tissue to secrete oestrogens. In the early stages, germ cell tumours of the testis may be impalpable, so estimations of serum hCG are important, particularly since chorionic carcinoma is the most common testicular tumour to cause gynaecomastia. Ultrasound of the testis can detect the presence of a small tumour.

Leydig cell tumours are rare before puberty, but they should be considered as a cause of gynaecomastia in adult males. Seminomas may cause gynaecomastia, but this is rare. The rarest testicular tumour of all, the Sertoli cell tumour, frequently presents with gynaecomastia and loss of libido.

**Thyroid disorders**

Eighty per cent of males with thyrotoxicosis have histological evidence of gynaecomastia. Plasma oestradiol levels are elevated. This is thought to be due to increased androstenedione production and increased oestrogen production in extraglandular sites. Increased oestrogen levels and excess thyroid hormones both lead to a rise in SHBG levels, so that more testosterone is bound and the ratio of free testosterone to free oestradiol falls.

Gynaecomastia occurs rarely in patients with hypothyroidism; the mechanism is uncertain.
Adrenal gland tumours
Adrenocortical carcinoma and, more rarely, adenoma may produce oestrogens and lead to the development of gynaecomastia. However, most suprarenal tumours produce large amounts of adrenal androgens (e.g. androstenedione and dehydroepiandrosterone). These androgens are metabolized by aromatization (aromatase enzyme), producing large amounts of oestrogen. Plasma or urinary oestrogen levels are therefore raised, and gonadotrophin levels may be suppressed. Urinary 17-oxo-steroid levels can be increased or normal. The testes are small, and aspermia is present. Testicular biopsy may show hypoplasia of Leydig cells. Computed tomography scanning of the adrenal gland should be carried out to localize the tumour.

Similarly, in congenital adrenal hyperplasia, the increased oestrogen production is due to increased production of androstenedione, which is metabolized by extraglandular aromatase to oestrogen.

Pituitary tumours
Growth hormone-secreting tumours (somatotroph adenoma) of the pituitary causing acromegaly, prolactinoma, or non-functioning pituitary adenoma (gonadotrophin staining) not producing excess growth hormone, may result in the production of prolactin (Fig. G.33). However, prolactin does not promote growth or development of the breast tissue and is therefore believed not to play a direct role in the development of gynaecomastia.

If gynaecomastia develops in association with a prolactin-secreting tumour, it is thought to be the result of secondary hypo-gonadism and an alteration in the ratio of free testosterone to free oestradiol. Some pituitary tumours have caused gynaecomastia because of the secretion of luteinizing hormone.

Hypothalamic disorders
Lesions in the hypothalamus may give rise to precocious puberty and gynaecomastia.

Liver dysfunction
In chronic liver disease (e.g. cirrhosis), there is a decline in testosterone production. In addition, there is a decreased clearance of androstenedione by the liver, and therefore more is available for extraglandular aromatization to oestrogen. The SHBG levels rise, and this further reduces the level of free testosterone. The resultant decrease in the androgen: oestrogen ratio produces the feminization and gynaecomastia associated with chronic liver disease.

Nutrition
Re-feeding after a period of starvation or malnutrition is associated with gynaecomastia that eventually regresses once nutrition is maintained. The mechanism of feminization is not well understood, but it may have a similar pathogenesis to that associated with chronic liver disease – that is, decreased clearance of androgens by the liver, making them available for extraglandular aromatization to oestrogen.

Another interesting theory regarding poor nutrition was noted in starving ex-prisoners of war when they were re-fed. Gonadotrophin levels, which are depressed during the period of starvation, rise following the receipt of food. The testes become stimulated again and the individual goes through what is, in effect, a second puberty.

Respiratory disorders
Carcinoma of the bronchus may secrete hCG, which, by stimulating the testes, leads to increased oestrogen production. In most cases where bronchial carcinoma has caused gynaecomastia, there has been associated hypertrophic pulmonary osteoarthropathy.

Drugs
Drugs are an important cause of gynaecomastia, especially in adults. Oestrogen therapy invariably produces gynaecomastia, and when diethylstilbestrol is used, a deep brown pigmentation of the nipple and areola develops (see Fig. G.31). Gynaecomastia has been described in pharmaceutical industry workers
involved in the manufacture of oestrogens. In particular, young men and boys are very sensitive to the effects of oestrogen. Exposure may occur through contact with oestrogen-containing creams, or oestrogen present in meat and dairy products from oestrogen-treated animals. hCG administration may cause gynaecomastia when used for the treatment of undescended testes. Methyltestosterone can occasionally lead to breast enlargement, possibly because of peripheral conversion to oestrogens. Any drug that is an androgen antagonist (see Box G.4) may cause gynaecomastia by allowing the unopposed action of oestrogens on breast tissue. Digoxin is associated with gynaecomastia in approximately 10 per cent of treated men, although the underlying mechanism is not fully understood. Although digoxin does not cause a rise in plasma oestrogen, it does bind to the human oestrogen receptor, and may promote gynaecomastia by enhancing the action of endogenous oestrogens. Antiretroviral therapy for HIV infection (particularly with the drug efavirenz) has been associated with gynaecomastia, although the mechanism is currently obscure. Clomiphene acts as an anti-oestrogen by blocking oestrogen receptors and is used to treat gynaecomastia in boys. It induces gonadotrophin release by interfering with the negative feedback effects of oestrogen on the hypothalamus, and therefore it may cause gynaecomastia on discontinuation of treatment due to luteinizning hormone effects on oestrogen production by the testes. Any drugs that interfere with the synthesis of testosterone by the testes for sufficient time will produce feminization and gynaecomastia by lowering the androgen: oestrogen ratio. Examples include drugs from the imidazole group (e.g. ketoconazole, metronidazole and etomidate), which interfere with steroid hormone synthesis in Leydig cells, and antineoplastic drugs through long-lasting toxic effects on Leydig cells (e.g. alkylating agents used for systemic or testicular neoplasms).

Spironolactone affects testosterone activity by interfering with testosterone synthesis (high dose) and blocking androgen receptors (low dose). Gynaecomastia is found in 50 per cent of men who receive 150 mg spironolactone per day. When the active metabolite canrenoate is used instead of the parent drug, gynaecomastia does not develop. Anti-androgens, by inhibiting testosterone binding to the androgen receptor, cause gynaecomastia. Examples of anti-androgens include cyproterone acetate, flutamide, zanoterone and bicalutamide. Cimetidine, and far less commonly ranitidine, causes gynaecomastia by blocking the androgen receptor. Cimetidine may also prevent the breakdown of oestradiol.
CAUSES OF HAEMATEMESIS

Gastrointestinal haemorrhage commonly presents as vomiting of blood or bloodstained gastric content; this is defined as haematemesis. Causes of haematemesis are listed in Box H.1.

The most common causes of profuse haematemesis are acute gastric or oesophageal erosions, gastric- or duodenal ulcer and cirrhosis of the liver. A long history of dyspepsia may be present in patients bleeding from gastric or duodenal ulcer, but this may not always be present, particularly in the elderly. Acute erosions are particularly common in patients taking aspirin or other non-steroidal anti-inflammatory drugs (NSAIDs), or they may follow acute alcohol ingestion. A history of alcoholism may point to cirrhosis of the liver. Endoscopic examination of the oesophagus, stomach and duodenum is essential in all cases of haematemesis.

SWALLOWED BLOOD

Bleeding from the nose (epistaxis), gums (scurvy, haematological malignancies), throat, lungs (haemoptysis) may be followed by haematemesis when blood has been swallowed and then vomited; this may occur at night when blood has been swallowed during sleep.

Spurious

Some patients suffering from a variant of the Munchausen’s syndrome may swallow blood in secret, and subsequently vomit this with an intent to deceive.

DISEASES OF THE OESOPHAGUS

Hiatus hernia and reflux oesophagitis

Hiatus hernia is common but is mostly associated with the symptoms of acid reflux rather than bleeding. Blood loss in oesophagitis and hiatus hernia is usually chronic, presenting as iron-deficiency anaemia, but haematemesis can occur.

Oesophageal varices

These are dilated veins, in the distal oesophagus (Fig. H.1) or the upper stomach as a result of portal hypertension due to cirrhosis of the liver. They can cause profuse haematemesis if rupture occurs. Stigmata of chronic liver disease may be present, such as jaundice, ascites, palmar erythema or spider naevi. The absence of these features does not exclude the presence of portal hypertension and varices.

Mallory–Weiss syndrome

A small tear of the mucosa at the oesophagogastric junction may result in haematemesis. The relatively common clinical picture is of retching and vomiting followed by haematemesis.

Oesophageal ulcer

Such ulcers may be either benign or malignant, and they may be associated with hiatus hernia and reflux oesophagitis. Although a relatively uncommon cause...
of haematemesis, significant bleeding may occur, the diagnosis being confirmed by endoscopy with biopsy.

**Mediastinal tumour penetrating the oesophagus and aorta**
This is an infrequent complication of such tumours, but it may occur if the tumour erodes into the oesophagus. It may be associated with compression and invasion of the large veins, leading to oedema of the neck and extremities, cyanosis and dilated superficial veins.

**Foreign body perforating oesophagus and aorta**
This rare cause of haematemesis may be induced by a fish bone, pin or dental plate perforating both the oesophagus and a large vessel or aorta. A history of a foreign body being swallowed, followed by a feeling of discomfort in the oesophagus, suggests the condition, which is generally confirmed by radiology or endoscopy.

**DISEASES OF THE STOMACH**

**Gastric ulcer**
Haematemesis may occur in acute or chronic gastric ulcer. A gradual loss of blood may allow sufficient time for gastric acid to convert haemoglobin into haematin, which gives the vomit a dark brown or ‘coffee-grounds’ appearance. Severe bleeding may occur if a medium-sized or large vessel is eroded. Profuse haemorrhage causes a feeling of faintness, restlessness, syncope and a rapid feeble pulse. There may be abdominal pain, nausea, vomiting and associated melaena. The pain is generally epigastric, but many haematemeses from ulcers are associated with no abdominal discomfort. An endoscopy is the most important initial procedure. The source of bleeding can be identified and treated using different endoscopic techniques. All gastric ulcers must be biopsied to exclude malignancy and the endoscopy repeated after 8 weeks to confirm that the ulcer has healed.

**Acute gastritis and haemorrhagic erosions**
In acute gastritis, the mucosa is congested, and small haemorrhages and erosions are identified at endoscopy. Erosions are small or minute ulcers, the differences between these and multiple small gastric ulcers being that of degree rather than of kind.

Gastritis and gastric ulcers are most commonly caused by the organism *Helicobacter pylori*. The other important cause are NSAIDs/aspirin. Alcohol and other medication (including bisphosphonates and clopidogrel) are also important to consider.

Gastritis and haematemesis may also be due to corrosive poisons, strong acids or alkalis destroying the surface membranes of mouth, throat, oesophagus or stomach, causing intense pain, dysphagia, retrosternal discomfort, abdominal distension, collapse and haematemesis. In arsenic poisoning, the mucous membrane of the stomach is red, inflamed, partly detached and covered with bloodstained mucus. The principal symptoms are nausea, severe sickness, burning epigastric pain and diarrhoea. The vomitus is usually a brown, turbid fluid mixed with mucus and streaked with blood in which arsenic may be detected by appropriate tests; severe diarrhoea may come later.

**Tumours**
Severe haematemesis is relatively rare in carcinoma of the stomach, accounting for less than 5 per cent of all cases in Western countries. The patient may have epigastric discomfort, nausea, vomiting, anorexia and weight loss. Pyrexia, anaemia, cachexia and an abdominal mass is present in advanced cases. Sometimes, no preceding symptoms occur, and the patient presents with haematemesis. If narrowing at the pyloric area has occurred, the patient may have regular vomiting streaked with blood of ‘coffee-grounds’ appearance. If the tumour is at the cardia, regurgitation of food immediately after eating may occur rather than true vomiting Troisier’s sign – enlargement of the left supra-clavicular lymph node – is a rare finding, but if present it indicates an intra-abdominal malignancy.
Gastric tumours are best diagnosed by endoscopy, through which confirmatory biopsies can be obtained. Gastrointestinal stromal tumours (GISTs) are less common, slow-growing submucosal tumours arising from the muscularis. They are often seen coincidentally at endoscopy but can ulcerate and bleed.

**Pseudoxanthoma elasticum (Grondblad–Strandberg syndrome)**

This is a hereditary disease characterized by widespread atrophy of elastic tissue throughout the body. Disintegration of the submucosal artery elatica leads to severe haemorrhage from the stomach and is a rare cause of haematemesis. Characteristic appearances of the skin of head, neck and body may suggest the diagnosis; these have been termed the ‘plucked chicken’ appearance. Angioid breaks in the retina are also seen; these can lead to visual loss (Fig. H.2).

**Hereditary haemorrhagic telangiectasia (Osler–Rendu–Weber syndrome)**

This is caused by a focal dilatation of postcapillary venules which have excessive layers of smooth muscle devoid of elastin. There are multiple telangiectases, which are commonly found on the lips and mucous membranes of the mouth and nose, and throughout the gastrointestinal tract. There may also be arteriovenous fistulae in the lungs, liver and brain. The condition is inherited in an autosomal dominant manner. The most common presentations are gastrointestinal tract bleeding, often haematemesis, sometimes melaena, or epistaxis. Endoscopic examination of the stomach generally identifies such lesions, but milder bleeding distal to the stomach is often difficult to diagnose.

**DISEASES OF THE DUODENUM**

**Duodenal ulcer**

These are caused by either *Helicobacter pylori* infection or NSAIDs/aspirin. Haematemesis occurs when a duodenal ulcer erodes a blood vessel, some of the blood regurgitating through the pylorus into the stomach to be vomited, the rest passing down the gastrointestinal tract to cause melaena. If bleeding is severe, red blood can occasionally be passed per rectum. There may be a dyspeptic history with epigastric pain occurring some hours after food, often at night, with symptoms showing periodicity. Generally, endoscopy can rapidly identify the presence and site of a bleeding duodenal ulcer, and enable treatment. Biopsies are also taken from the stomach to detect *Helicobacter pylori* infection.

**Tumours**

Duodenal tumours are rare, mostly arising from the ampulla of Vater, and associated with inherited polyposis syndromes (familial adenomatous polyposis). They may be missed at endoscopy.

**Gallstones ulcerating into the duodenum**

This is a rare cause of haematemesis and is generally associated with previous attacks of colicky abdominal pain, classically under the right costal margin, and sometimes associated with jaundice. The diagnosis is generally confirmed by plain abdominal X-ray showing air in the biliary tree, ultrasound examination, the passage of stone in the faeces, or endoscopy.

**PORTAL HYPERTENSION**

**Chronic liver disease**

Chronic liver disease is associated with portal hypertension (oesophageal and gastric varices) as well as with disordered haemostasis from thrombocytopenia from portal hypertension, and from the impaired production of hepatic clotting factors.

**Portal vein obstruction**

Extrahepatic portal vein thrombosis usually occurs in infants and early childhood. The cause is usually unknown, and infection is rare. The patient presents with haematemesis from oesophageal varices, or splenomegaly. There is no evidence of liver dysfunction clinically or biochemically. The outlook is good. Surgery should be avoided, but if bleeding is a problem, some form of portal systemic shunt surgery is required.

In adults, portal vein thrombosis is usually due to a procoagulant state (either acquired due to medication such as the oral contraceptive pill, or inherited) or due to trauma, surgery or neoplastic invasion.
DRUGS
A number of drugs may be associated with haematemesis, the cause being suspected by an accurate and detailed drug history from the patient, or the presence of a disorder such as arthropathy, suggesting that the patient may be having – or has had – drugs such as NSAIDs that may lead to gastrointestinal haemorrhage.

Anticoagulant and antiplatelet therapy
Anticoagulants may be indicated in various conditions, including deep venous thrombosis or pulmonary thromboembolism, and patients on long-term anticoagulant therapy may have significant gastrointestinal haemorrhage from relatively minor lesions in the stomach or duodenum, such as small ulcers or erosions. Antiplatelet medications used in a variety of vascular diseases, in particular the acute coronary syndromes, increase the risk of gastrointestinal haemorrhage. These include well-established drugs such as aspirin and dipyridamole. Newer drugs now being more commonly prescribed include clopidogrel, and new oral anticoagulants e.g. rivaroxaban, apixaban, dabigatran.

NSAIDs
These drugs are used in all forms of arthropathy, and they may lead to the development of haemorrhagic erosions or erosive gastritis in the stomach or duodenum, with haematemesis. These are in the stomach or duodenum, with haematemesis. They are widely used, especially in elderly patients. An accurate drug history should provide the clue to the cause, but endoscopy will localize the site and assess the severity of the lesion.

MISCELLANEOUS
A number of rarer conditions may be associated with gastrointestinal haemorrhage and haematemesis. These include: severe bleeding from an abdominal aneurysm opening into the duodenum (aorto-duodenal fistula); uraemia (in which the presence of high blood pressure, cardiac hypertrophy,); abdominal surgery; trauma or burns (Curling’s ulcer); and autoimmune disorders such as polyarteritis nodosa or systemic lupus erythematosus.

HAEMATOMSPERMA
Ben Challacombe
The alarming symptom of passing blood in the ejaculate is extremely worrying and distressing to most men. Thankfully, haematospermia or haemospermia, although of sudden onset, is rarely a sign of significant pathology and is usually painless and self-limiting, with no sinister cause being found. The true incidence of haematospermia is probably unknown as most ejaculates go unnoticed during sexual intercourse. In younger men under the age of 40, it is commonly due to inflammation from prostatitis, urethritis and epididymo-orchitis. Less commonly, bleeding is related to trauma to the perineum or testes, and it can more rarely be related to testicular tumours. Following clinical examination, this group of men can generally be reassured and counselled regarding the likely time period for complete resolution (2-4 weeks). Clearing of the ejaculate is related to the number and frequency of ejaculations, and patients should be informed that the semen will often change from bright red to darker red and finally to brown in colour before returning to normality.

In older men, the above causes are also seen, but another group of significant pathologies is also possible. These include prostate cancer, bladder cancer, benign prostatic hyperplasia, dilated prostatic veins, prostatic or seminal vesicle stones, severe hypertension and rare cancers of the seminal vesicles. Haemospermia is now commonly seen after transrectal or trans-perineal ultrasound-guided prostate biopsy as part of the investigation for prostate cancer, where it is usually transient and self-limiting.

Rarer causes include bleeding diatheses, tuberculosis, congenital abnormalities (Müllerian and urticarial cysts), schistosomiasis, hydatid disease, HIV and amyloid.

A thorough clinical examination is required and should include the abdominal, rectal and genital regions (including palpation of the vas/spermatic cord), as well as a blood pressure assessment. Primary investigations include urine dipstick, prostate-specific antigen (in patients over 40 years of age) and semen culture. The main initial aim is to exclude significant underlying pathology, and to reassure the patient. If associated with haematuria, then cystoscopy and renal tract imaging should also be completed (see HAEMATURIA, below).

In persistent or recurrent haematospermia, transrectal ultrasound should be considered along with urine cytology, flexible/rigid cystoscopy and dynamic magnetic resonance imaging (MRI) or computed tomography scan of the pelvis.

HAEMATURIA
Ben Challacombe
Haematuria means the presence of red blood cells in the urine, and although free haemoglobin may be present in the urine as a result of lysis of cells in the urinary tract, it should not be confused with haemoglobinuria, in which the pigment alone is filtered
through the glomeruli. In clinical practice, there are two main ways in which haematuria may pose a diagnostic problem. Macroscopic (gross/visible) haematuria may be a presenting feature, with or without other symptoms, or blood may be found in the urine only by ‘dipstick’ testing or microscopy. Microscopic haematuria is defined as the presence of three to five red blood cells per high-power field on microscopy. Both macroscopic and microscopic haematuria require investigation, but the chance of finding a urinary tract malignancy (up to 30 per cent have bladder or renal tumours) or other significant pathology (stone or bleeding in benign prostatic hypertrophy) is far more common in the former. In microscopic haematuria, many cases are related to medical problems such as renal disease, diabetes and hypertension, although urinary tract malignancy always needs exclusion. Referral to an urologist is recommended for the investigation of haematuria.

There are a number of false-positive diagnoses for haematuria, including menstruation in women, after vigorous exercise, after sexual intercourse, in the presence of bacterial peroxidases, with povidone and with myoglobinuria. Frank haematuria (macroscopic or visible haematuria) may be painful or painless. The colour of the blood may be of some significance (although this is often unreliable) as, if it is bright red, the blood is more likely to have come from the lower urinary tract (bladder or urethra). Initial haematuria, when bleeding occurs at the start of micturition, implies that the bleeding may be coming from the urethra, prostate or bladder neck area. Blood that is mixed with the urinary stream may have a source in the kidneys, ureters or bladder. The amount of blood present is also of diagnostic importance; in the absence of trauma, a large quantity of blood is suggestive of a tumour of the urinary tract, although profuse bleeding is quite common in several other conditions, for example benign prostatic hypertrophy. Painless haematuria is more likely to be due to neoplasia – bladder tumours most commonly present with this symptom. Painful haematuria is more common where there is associated bladder infection or calculi, but tumours cannot be excluded on the basis of clinical history alone.

The history should include an enquiry into the urological symptoms, along with the patient’s past and family history, together with details of his or her occupation (including exposure to chemicals and dyes), smoking history and any drugs being taken. Loin pain may be associated with a renal lesion and can occur in renal tumours and obstruction. Loin pain that radiates to the groin and is colicky in nature suggests renal colic due to the passage of a stone or blood clot. Increased urinary frequency or penile pain immediately after micturition suggests bladder disease. Pain in the pelvic or sacral area with haematuria suggests malignant disease in the bladder or prostate. The site of associated symptoms may, however, be misleading. For example, a tumour in the bladder may, by occluding the ureter, cause unilateral hydronephrosis with loin pain, or tuberculosis of the kidney and ureter can cause increased frequency of micturition in the absence of bladder infection.

Physical examination in a patient with haematuria is usually unremarkable, but significant anaemia should be looked for. There are no specific signs of haematuria except blood at the external meatus, which may be seen in association with urethral trauma (as a result of catheterization or pelvic fracture injury). The kidneys should be palpated to determine their size, if possible, and to elicit any tenderness. Kidneys are not normally palpable except in very thin subjects, and palpation requires experience. Suprapubic palpation may detect a distended bladder due to clot retention. A palpable bladder mass is rare except when large tumours are present. Rectal and vaginal examination will allow palpation of the pelvic viscera. On rectal examination, the uniform, elastic and movable prostate of benign prostatic hyperplasia can be distinguished from the hard, nodular, often immovable gland without a median groove characteristic of prostatic carcinoma. Lymphatic spread from a carcinoma of the bladder or prostate may be palpable as thickening in the lateral pelvic space. Vaginal examination will also allow palpation of the bladder base and lateral pelvic space, as well as of the other pelvic organs; in the fornices, the lower end of each ureter can sometimes be felt if it is diseased or contains a calculus. The testes should be examined, particularly for evidence of tuberculosis of the epididymis.

Macroscopic inspection may reveal clots. Their shape may suggest the source of bleeding; clots formed in a renal pelvis may be triangular in shape, while those formed in a ureter are likely to be thin and ‘worm-like’. The urine itself should be examined both by dipstick analysis and by sending a mid-stream urine (MSU) sample for microscopy and culture. The presence of red-cell casts is pathognomonic of glomerular bleeding. Large numbers of oxalate crystals may indicate a tendency to oxalate stone formation; the much rarer crystals of cystine are diagnostic of cystinuria, with its strong tendency to the formation of calculi. The finding of clumps of transitional epithelial cells is suggestive of carcinoma of the bladder, and formal cytological examination of the urine should be organized. Urine cytology or one of the newer urinary tests for bladder cancer (NMP22 or BT-STAT)
is part of the standard haematuria investigation in many centres. Rarely, fragments of renal papillae may be seen, sometimes with the naked eye, in papillary necrosis associated with chronic interstitial nephritis. The presence of leucocytes is not very helpful as they are likely to be found not only in bacterial infections but also in tuberculosis, tumours and benign prostatic hypertrophy.

Further investigation of haematuria should include:

- Cystoscopy: flexible or rigid depending on the severity of bleeding and patient choice.
- Upper urinary tract imaging: investigation policies differ between hospitals
  - Increasingly a contrast plus non-contrast computed tomography (CT) scan is being used as the primary investigation of haematuria. A CT scan of the abdomen and pelvis is also necessary when significant abnormalities are found on ultrasonography or in the staging of renal and bladder cancer.
  - A kidney–ureter–bladder X-ray and ultrasound of the urinary tract may be performed.
  - Intravenous urography (IVU) is less commonly performed nowadays as ultrasonography produces useful information and will pick up urinary tract tumours in bladder and kidneys. Where transitional cell carcinoma (TCC) or ureteric calculus is suspected, an IVU should be performed.

The main causes of haematuria are summarized in Box H.2.

**Box H.2 Causes of haematuria**

<table>
<thead>
<tr>
<th>Renal</th>
<th>Vesical</th>
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<tbody>
<tr>
<td>Trauma</td>
<td>Trauma</td>
</tr>
<tr>
<td>Tumours</td>
<td>Tumours</td>
</tr>
<tr>
<td>Calculus</td>
<td>Prostatic enlargement</td>
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<td>Glomerulonephritis</td>
<td>Tuberculosis</td>
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<td>Polycystic kidneys</td>
<td>Calculus</td>
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<tr>
<td>Tuberculosis</td>
<td>Cystitis</td>
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<tr>
<td>Pyelonephritis</td>
<td>Foreign body</td>
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<tr>
<td>Infarction</td>
<td>Disease of adjacent organs</td>
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<tr>
<td>Polycystis nodosa</td>
<td></td>
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<tr>
<td>Chronic interstitial nephritis</td>
<td></td>
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<tr>
<td>Hydronephrosis</td>
<td></td>
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<tr>
<td>Irradiation nephritis</td>
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<tr>
<td>Hydatid disease</td>
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<tr>
<td>Medullary sponge kidney</td>
<td></td>
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<tr>
<td>Relief of tension</td>
<td></td>
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<tr>
<td>Ureteric</td>
<td>Urethral</td>
</tr>
<tr>
<td>Calculus</td>
<td>Urethritis</td>
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<tr>
<td>Tumours</td>
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<tr>
<td>General</td>
<td>Caruncle</td>
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<td>General</td>
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<tr>
<td></td>
<td>Drugs</td>
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<td></td>
<td>Bleeding disorders</td>
</tr>
</tbody>
</table>

**RENAL CAUSES OF HAEMATURIA**

Haematuria can follow trauma of any degree of severity. A history of an accident or a blow or kick to the lumbar region that is associated with haematuria suggests damage to the kidneys. Even slight injury to the loin, of which there may be no recollection or external sign, can cause haematuria, especially if there is a pre-existing renal lesion (horseshoe kidney, tumour or polycystic kidney). The kidney may be palpable, but this may be due to extravasation of blood or urine forming a perinephric haematoma/urinoma.

Trauma to the urinary tract may give rise to haematuria, and the site of injury usually indicates the likely source of bleeding. Thus, pelvic fracture may be accompanied by haematuria due to urethral injury. Urethral injury may be contusion or rupture. Blood at the external meatus, although signifying a likely urethral injury, does not mean that the urethra has been completely ruptured. Thus, the gentle passage of a urethral catheter under aseptic conditions may be tried without causing further damage. Traumatic bladder injury may give rise to extravasation. However, the signs may be masked by the associated pelvic injury. A CT scan will help in making the diagnosis, and is recommended in cases of macroscopic haematuria following trauma.

Renal tumours are important causes of haematuria. Although today many tumours are found incidentally, the most common presenting symptom in carcinoma of the kidney is profuse intermittent painless haematuria. A mass may occasionally be felt in the loin, and there may be pain in that region, resulting from increasing tension or colic from the passage of clots. Unexplained fever, polycythaemia and hypercalcaemia may be present (paraneoplastic features), and hypertension is present in about 30 per cent of cases. An ultrasound scan will demonstrate a mass with mixed echogenicity. When a tumour is suspected on ultrasound scanning, a contrast-enhanced triple phase or renal protocol CT scan will confirm the diagnosis. Renal arteriography is not usually performed in the investigation of renal tumours except when a partial nephrectomy (renal conservation) is proposed, so that the vascular anatomy of the kidney can be studied prior to surgery.

In children with nephroblastoma (Wilms’ tumour) the presenting feature is commonly abdominal distension and a mass is almost always palpable; haematuria occurs later.

An angiomyolipoma of the kidney may present with visible bleeding, but is more likely to present...
with pain in which bleeding has occurred into the perinephric tissues. CT scanning is necessary to diagnose it specifically, with the fat component, showing black on the images, usually confirming the diagnosis (Fig. H.3).

Transitional cell carcinoma of the renal pelvis is uncommon, but it may cause profuse haematuria that is usually painless but can be associated with clot colic. Ultrasonography may show a filling defect in the renal pelvis (Fig. H.4) and CT-IVP can confirm the diagnosis.

Renal calculus seldom causes profuse bleeding. Haematuria (microscopic or occasionally macroscopic) and often pyuria may occur. An ache in the loin is common while the stone remains in the kidney, and it may be followed by renal colic if a smaller stone passes down the ureter. This typical, very severe pain passes from the loin downwards and forwards to the groin, upper part of the thigh and testicle, and is accompanied by urinary frequency and urgency (strangury). The radiographic diagnosis of ureteric calculus is discussed on p. 246. A renal calculus may become too large to pass into the ureter, and it may then cause hydronephrosis, pyonephrosis, sepsis and renal failure, with corresponding symptoms that may include haematuria (Fig. H.5).

Nephrological disease is a common cause of microscopic and occasionally macroscopic haematuria. An important cause of recurrent macroscopic or microscopic haematuria is Berger’s nephritis (IgA nephropathy), which is the most common glomerulonephritis throughout the world. The episodes of haematuria commonly follow a few weeks after upper respiratory tract infections; the characteristic histological feature is the deposition of IgA in the glomeruli. The condition mainly affects children and young adults and, although the prognosis is good in most, there is a slow progression to chronic renal failure in 25–30 per cent of cases over a period of 20 years. Glomerular IgA deposition is also a common feature of Henoch–Schönlein syndrome, which, although occasionally severe, is more often a relatively benign condition characterized by haematuria with typical purpuric and oedematous skin lesions, arthritis and abdominal pain with intestinal bleeding. A renal biopsy is needed to confirm diagnosis.
Post-streptococcal glomerulonephritis is now rare in the Western world, but it is still common in developing countries, where it often follows streptococcal pharyngitis or skin infections. The characteristic features are macroscopic haematuria, producing the typical ‘smoky’ appearance, proteinuria, oliguria, generalized oedema and hypertension. A similar lesion is a common complication of infective endocarditis, in which the glomerular changes are focal and segmental. Examination of a fresh specimen of urine will show red cells in a majority of cases of endocarditis and is thus an important diagnostic procedure. In both these conditions, the glomerulitis is due to the deposition of immune complexes.

Rapidly progressive glomerulonephritis is a cause of macroscopic haematuria with loin pain; it causes rapid deterioration and can lead to acute renal failure within a few days. It can occur in Henoch–Schönlein syndrome, cryoglobulinaemia, microscopic polyarteritis, Wegener’s granulomatosis and the nephritis associated with the development of antibodies to glomerular basement membrane (anti-GBM nephritis). The latter is often associated with pulmonary haemorrhage (Goodpasture’s syndrome).

A number of hereditary varieties of nephritis can also cause haematuria; of these, the most common is Alport’s syndrome, in which a renal lesion, often severe but less so in females, is associated with nerve deafness; the differential diagnosis, which can be made by renal biopsy, includes familial benign haematuria, in which proteinuria and deafness do not occur, and intermittent or persistent microscopic haematuria continues for many years without any deterioration in renal function.

Nephritis with haematuria is also a feature of several systemic diseases, of which the most important is systemic lupus erythematosus (SLE); of the numerous manifestations of this disease, arthritis, cutaneous lesions and hypertension are those most often associated with nephritis. Haematuria is not one of the typical features of the nephrotic syndrome, but some of the causes of the latter, including SLE, produce sufficient glomerular inflammation to cause haemorrhage. Apart from this, renal vein thrombosis, which is a recognized complication of the nephrotic syndrome, causes a sudden deterioration in renal function together with haematuria, which is often macroscopic. Other important associations with chronic renal disease that may cause microscopic haematuria are atherosclerosis, hypertension and diabetes.

Adult polycystic disease of the kidneys is frequently associated with haematuria, either painless or painful, due to clot colic or an acute bleed into a cyst; it can be precipitated by mild trauma. Both kidneys are usually palpable, and hypertension is common. The differential diagnosis includes bilateral hydronephrosis due to a lesion of the bladder, prostate or urethra causing obstruction and often haematuria. The CT and IVU appearances are quite different: in polycystic disease, the calyces are narrow and elongated, quite unlike the dilated pelvis in hydronephrosis.

Haematuria in renal tuberculosis is usually slight and is associated with pyuria; occasionally, an episode of gross haematuria is the presenting feature. Currently, tuberculosis predominantly affects Asian populations, with a higher incidence in males, and those with HIV/AIDS. Haematogenous spread from the lungs leads to granuloma formation in the renal cortex. Patients may complain of a dull ache in one loin, with occasional exacerbations resembling renal colic. Once the tuberculous focus has ruptured into the renal pelvis, the characteristic symptom is increased frequency of micturition both by day and by night, even without any involvement of the bladder. The thickened lower end of the ureter may be palpable on rectal or vaginal examination and, in males, nodules may be felt in the prostate or seminal vesicles. The cystoscopic appearance of the ureteric orifice may be distinctive.
HAEMATURIA

Caseous necrosis of the renal papillae and deformity of the calyces leads to the release of organisms into the urine. Hydronephrosis may develop in the affected kidney, and IVU or CT scanning will reveal characteristic changes. Eventually, there is fibrosis and calcification, leading to renal destruction and auto-nephrectomy. The diagnosis can be confirmed by special bacteriological culture of early morning specimens of urine. Other infections of the kidney, such as acute pyelonephritis, occasionally cause haematuria, which is seen more commonly in women.

Vascular lesions of the kidneys can cause haematuria. Infarction is usually due to an embolus arising, for example, from a fibrillating left atrium, mural thrombus following myocardial infarction, a large vegetation in infective endocarditis or, very rarely, left atrial myxoma. Infarction can also occur in macroscopic polyarteritis nodosa. Sickle-cell disease may cause haematuria as a consequence of vascular damage, and aneurysm of the renal artery and intrarenal arteriovenous fistula are also rare causes of haematuria. Accelerated (‘malignant’) hypertension can cause haematuria, which is usually microscopic but occasionally macroscopic.

Less common causes of renal origin haematuria include loin pain/haematuria syndrome, which is a diagnosis of exclusion, and medullary sponge kidney, which is usually asymptomatic, although haematuria and renal colic may occur as a result of calculus formation.

URETERIC CAUSES OF HAEMATURIA

Ninety per cent of ureteric calculi will be associated with haematuria, which is usually picked up on urine dipstick examination or microscopy and is usually (90 per cent) microscopic.

Transitional cell carcinoma of the ureter is rare; it can be diagnosed by finding a filling defect in the ureter with dilatation above on a CT–intravenous pyelogram, IVU or retrograde ureterography or ultrasound scan showing a dilated ureter. The differential diagnosis of the negative shadow produced by a tumour in the ureterogram includes a radiolucent calculus and an air bubble.

Other ureteric causes include trauma, most often due to iatrogenic injury during laparotomy, and ureteric strictures.

VESICAL CAUSES OF HAEMATURIA

Intermittent profuse painless haematuria is characteristic of TCC or other tumours of the bladder, and indicates this diagnosis until proven otherwise. Other symptoms of bladder TCC or carcinoma-in-situ include urinary frequency, urgency and dysuria. Cystoscopy (flexible or rigid) is the essential diagnostic procedure. Bladder tumours may occur at any age, but they are rare below the age of 40 and most commonly occur in the 60s and 70s in current or prior smokers. Tumours may be single or multiple. Adenocarcinoma and squamous cell carcinoma of the bladder are rarer tumours associated with urachal lesions and chronic inflammation from stones or infections, respectively.

Profuse haematuria may occur in patients with benign or malignant prostatic enlargement. Vesical tuberculosis is seeded from the kidneys, and causes persistent frequency, slight haematuria and pyuria. Evidence of tuberculosis elsewhere, in the lungs or the genital organs, may be found.

A bladder calculus may produce haematuria due to direct trauma to the bladder neck from the stone or in association with urinary tract infection. Suprapubic pain usually accompanies this diagnosis.

Acute cystitis is often accompanied by haematuria; the diagnosis will usually be obvious on other grounds. There is, however, a form of haemorrhagic cystitis in which bleeding predominates over other symptoms; a similar form of haemorrhagic cystitis is a complication of treatment with cyclophosphamide.

Chronic interstitial cystitis more commonly in the female may be accompanied by painful haematuria and an increase in frequency of micturition due to reduced bladder capacity.

Radiation cystitis is characterized by multiple telangiectases in the vesical mucosa, which may bleed and prove extremely resistant to endoscopic treatment. Vesical schistosomiasis causes slight haematuria and other symptoms similar to those of vesical tuberculosis. There is likely to be a history of residence in an endemic area such as Egypt and neighbouring countries. The typical ova of Schistosoma haematobium may be found in the urine and, at cystoscopy, so-called ‘sandy patches’ may be seen near the trigone; these consist of numerous ova without any inflammatory reaction.

The sudden emptying of the bladder in a case of chronic urinary retention may cause decompression haematuria, which usually comes from vessels within the bladder wall or occasionally from the kidney. It is self-limiting; a slow release of urinary retention/slow decompression does not prevent this, and has no place in the modern management of chronic urinary retention.

Foreign bodies may be introduced into the bladder by accident or deliberately and cause bleeding; their presence will be revealed by cystoscopy or radiography. Haematuria may occur as a result of spread of disease.
of neighbouring viscera to the bladder. Carcinoma of the uterus, vagina, rectum or pelvic colon can all invade the bladder, usually at a late stage of the disease. Microscopic haematuria may result from the contact of an acutely inflamed appendix with the bladder wall and cause localized cystitis. This may be a source of diagnostic confusion, but the other symptoms of acute appendicitis are likely to be present. Rectal examination will reveal the inflammatory process in the right side of the pelvis. More rarely, acute salpingitis or pelvic abscess can cause haematuria by a similar mechanism. Haematuria can also be caused by a direct spread of inflammation from tuberculosis or dysenteric ulceration of the intestines, or from diverticulitis of the colon; the latter is particularly likely to lead to a vesicocolic fistula, with the classical additional symptoms of pneumaturia and recurrent infections.

URETHRAL CAUSES OF HAEMATURIA
Lesions of the urethra can cause blood to appear spontaneously at the meatus, as well as haematuria. Such conditions include acute urethritis, calculus and carcinoma. Foreign bodies may be introduced into the urethra as a form of sexual excitement and, in females, a urethral caruncle may develop at the urethral orifice and be visible on examination.

GENERAL CAUSES OF HAEMATURIA
Several drugs have been implicated as causes of haematuria, but anticoagulants are the only ones of practical importance. It must be remembered that the fact that a patient is on anticoagulants does not exclude other causes of haematuria, such as renal, bladder or prostatic carcinoma. Finally, thrombocytopenia and disorders of platelet function can cause haematuria, as can haemophilia, Christmas disease and, occasionally, scurvy. In these conditions, the diagnosis will usually be clear on other grounds.

HAEMOPTYSIS
Alex West

The coughing up of blood is called haemoptysis, a term generally accepted to refer specifically to expectoration of blood resulting from bleeding in the lungs or bronchi. The amount of expectorated blood may vary widely from slight streaking of the sputum to massive exsanguinating haemorrhage. Blood – either alone or mixed with sputum – is nearly always produced by coughing, but rarely may trickle past the larynx to ‘well up in the throat’. For this reason, some patients with true haemoptysis present to an otolaryngologist. Blood from the nasopharynx or larynx may lead to spitting of blood or bloodstained secretions. This is sometimes called spurious haemoptysis since the blood does not, in the strict sense, arise from the lungs.

In the assessment of haemoptysis, it is important to establish whether the bleeding has arisen from the chest, the nasopharynx or the upper gastrointestinal tract. Although examination of the nasopharynx should never be omitted, it is unusual to find bleeding lesions in the upper respiratory tract. Bleeding of oesophageal, gastric or duodenal origin can usually be differentiated from haemoptysis by the presence of gastrointestinal symptoms, such as nausea or vomiting, or a history of oesophageal varices or peptic ulcer disease. Prompt upper gastrointestinal tract endoscopy will settle the issue in doubtful cases.

Patients usually regard the presence of blood in the sputum as a sinister sign of serious lung disease. Haemoptysis may be a single event, and of little prognostic importance if a slight blood-staining of the sputum follows a repeated violent bout of coughing. An effortless haemoptysis of 1–2 ml of blood is much more likely to be of importance, especially if it is followed by the production of further bloodstained sputum. The history and physical examination, with special attention to the respiratory and cardiovascular systems (including the leg veins) and to the nasopharynx, may provide clues to the underlying cause of haemoptysis, but are seldom diagnostic. A chest X-ray is mandatory and may reveal evidence of infections including tuberculosis, old or new inflammatory lesions, possible malignancies or vascular abnormalities. Routine laboratory tests should include a full blood count and tests to exclude a bleeding diathesis. Sputum should be sent for microbiological testing both for microscopy, culture and sensitivity (MC&S) and for staining for acid fast bacilli. Sputum cytology may be of value but only if bronchoscopy is unavailable. If routine investigations are normal but haemoptysis persists, further imaging of the chest by computed tomography (CT) scanning is often helpful for example showing areas of localized bronchiectasis, endobronchial or central lesions not seen on the CXR, and to assist in identifying the presence of arteriovenous fistulae.

Haemoptysis, especially isolated instances of blood-staining of the sputum, often remains unexplained even after extensive investigations. Up to one-fifth of patients are in this category, and they should be followed up with further chest radiography after an interval of about 2 months, with the view also of obtaining the results of culture for mycobacteria and repeating investigations as necessary.

The causes of haemoptysis according to appearances on chest radiography are listed in Box H.3.
SPURIOUS HAEMOPTYSIS

Bleeding from the upper respiratory tract usually presents as obvious bleeding from the gums or from the nose (see EPISTAXIS, p. 162). As noted above, it may give rise to ‘haemoptysis’, because blood can be aspirated into the lungs during sleep and subsequently expectorated on awakening, possibly mixed with bronchial secretions. The nasopharynx should be examined in all cases of haemoptysis.

TRUE HAEMOPTYSIS

The patient’s age and environment are of importance. Where tuberculosis is a common disease, haemoptysis must always give rise to a suspicion of pulmonary tuberculosis, especially in younger patients. Bronchial carcinoma is a frequent cause of haemoptysis in middle-aged or older patients who smoke cigarettes. With either of these diseases, there may be a history of ill-health with non-specific respiratory symptoms preceding haemoptysis by a few weeks or months; however, the haemoptysis is often the first event to alert the patient to seek medical advice.

Pulmonary tuberculosis

Haemoptysis is often an early symptom of pulmonary tuberculosis. At this stage, physical examination is seldom abnormal, but radiographic changes are to be expected, with localized mottled shadowing or consolidation, possibly with cavitation. Rarely, haemoptysis may arise from radiographically obscure disease, with localized shadowing concealed by the overlying skeleton, hilar or mediastinal shadows. Lateral chest X-ray or CT scanning views may then be required for diagnosis. Haemoptysis in patients with active tuberculosis varies in severity from streaky staining of the sputum to profuse life-threatening bleeding. Severe haemoptysis may arise in chronic cavitated disease from rupture of an aneurysmal dilatation of an artery, exceptionally remaining patent in a strand of tissue traversing a cavity – the so-called ‘aneurysm of Rasmussen’. Old calcified tuberculous lesions may also be sufficient cause for haemoptysis simply due to local bronchiectasis, although reactivation of tuberculosis must be considered. Investigation of all cases in which there is radiological evidence suggesting active or inactive tuberculosis must include examination of the sputum for tubercle bacilli (acid-fast bacilli, AFBs).

Fungal infections

When more common diseases such as tuberculosis have been excluded, histoplasmosis, coccidioidomycosis and blastomycosis must be considered in the differential diagnosis of haemoptysis associated with abnormal appearances in the lung, especially in areas where these diseases are prevalent. Diagnosis depends upon isolation of the causal organism and may be aided by serological tests. Other rare infections that must be similarly considered when tuberculosis has been excluded include actinomycosis, nocardiosis and cryptococcosis; diagnosis of these depends upon isolation of the causal organism.

Bronchial carcinoma

Haemoptysis, an early symptom of bronchial carcinoma, usually takes the form of blood-streaking of sputum and possibly small free haemoptysis, often repeated over days or weeks. Later, more

Box H.3 Causes of haemoptysis on chest radiography

Radiological abnormality that is readily diagnosed
- Pulmonary tuberculosis
- Tumours of the lung – carcinoma, adenoma, etc.
- Pneumonia
- Pulmonary infarction
- Aspergillosis
- Contused lung due to trauma
- Mitral stenosis
- Large arteriovenous malformation

Radiological abnormality, the nature of which is not immediately obvious
- Pulmonary infarction
- Pulmonary haemosiderosis
  - Childhood haemosiderosis
  - Adult haemosiderosis
- Goodpasture’s syndrome
- Associated with systemic lupus erythematosus
- Associated with pulmonary vasculitis

With no gross radiological abnormality
- Mitral stenosis
- Pulmonary embolism
- Bronchitis and/or bronchiectasis
- Tumours of the larynx, trachea or larger bronchi not yet blocking the lumen
- Hereditary haemorrhagic telangiectasia and other pulmonary arteriovenous malformations

Bleeding diathesis

The primary disease is virtually always obvious from its other manifestations. Puzzling haemoptysis is rarely due to an unrecognized bleeding disorder

Iatrogenic
- Needle lung biopsy
- Transbronchial lung biopsy
severe bleeding may occur from the erosion of larger vessels, either by the tumour or by the suppuration, which often results from bacterial infection beyond it. Bronchial carcinoma must be suspected especially in cigarette smokers at or past middle-age, but it may occur in younger individuals. There is usually an obvious abnormality on the chest X-ray; the more common findings are of two sorts. The first abnormality is associated with tumours originating in large bronchi, and consists of air-absorption collapse or consolidation of a segment, a lobe (Fig. H.6) or even a whole lung beyond a complete obstruction, or patchy inflammatory consolidation beyond a partial obstruction. The second abnormality consists of localized (usually rounded) shadows in the lung fields, produced by tumours originating more peripherally. In some cases of squamous cell carcinoma, a rounded shadow of this sort may show a central transradiant area, due to necrosis of the central part of the tumour. Such appearances must lead to a provisional diagnosis of bronchial carcinoma. In occasional cases, a bronchial carcinoma arising in a large bronchus causes haemoptysis before it has obstructed the bronchus and before there is any abnormality on the ordinary chest X-ray taken in full inspiration. In such patients, there is likely to be a wheeze that may be mistaken for an asthmatic or bronchitic wheeze of expiratory airflow obstruction. Wheezes due to partial obstruction of larger airways differ in two respects. First, they may be localized and ‘fixed’ – that is, they do not clear on coughing. Second, they are as apparent in inspiration as in expiration – a fact that can be confirmed on spirometry; inspiratory flow rates are decreased as much as expiratory flow rates, resulting in characteristic changes in the volume–flow loops. Another way of recognizing partial obstruction of one of the lobar or main bronchi is to take a chest X-ray on full expiration. This may result in air-trapping in the affected lobe or lung. But ultimately patients with any suspected lung carcinoma should have a prompt CT scan where available.

Fibreoptic bronchoscopy is indicated to obtain histological specimens by brush biopsy, forceps biopsy and bronchial washings. In the case of peripheral tumours that cannot be reached by fibreoptic bronchoscopy, even under fluoroscopic control, transthoracic biopsy with a thin needle may be indicated. As transthoracic needle biopsy is a ‘rule in’ test (i.e. although a positive result is diagnostic, a negative result does not rule out carcinoma), patients should not be exposed unnecessarily to the risks of this when thoracotomy will be indicated, whatever the result. If available a PET scan is very helpful with diagnosis and staging.

**Bronchial adenoma or carcinoid tumours**

Haemoptysis is an important symptom in bronchial adenoma; these tumours may be highly vascular. Episodes of haemoptysis may occur over several years, and there may also be a history of recurrent attacks of pneumonia always involving the same lobe. There may be clinical and radiological evidence of lobar collapse or consolidation, possibly with abscess formation; an adenoma in a central bronchus may present with the manifestations of partial obstruction of a large airway, as described above for bronchial carcinoma. Since most of the tumours arise in the large bronchi, a bronchoscopic diagnosis is straightforward, but these tumours can bleed profusely on biopsy so caution is required.

**Pneumonia**

*Bacterial pneumonias* usually present as an acute illness with chest pain, dyspnoea, cough, fever and even rigors. The characteristic clinical features and the results of the blood count and direct examination of the Gram-stained smear of sputum that indicate the likely organism (for initiation of therapy), before the results of sputum cultures become available, are listed in Table H.1. Haemoptysis rarely amounts to more than blood-staining; the rusty sputum associated especially
HAEMOPTYSIS

Table H.1 Acute pneumonia in previously healthy adults and children

<table>
<thead>
<tr>
<th>Causes of pneumonia</th>
<th>Initial clinical features</th>
<th>Age</th>
<th>Blood count</th>
<th>Gram-stained sputum smear</th>
</tr>
</thead>
<tbody>
<tr>
<td>Streptococcus pneumoniae</td>
<td>Sudden onset with fever, cough and pleuritic chest pain</td>
<td>All ages especially adults</td>
<td>Leucocytosis &gt;12,000/mm³</td>
<td>Smear shows bacteria and large numbers of leucocytes</td>
</tr>
<tr>
<td>Staphylococcus aureus</td>
<td></td>
<td>Children</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haemophilus influenzae</td>
<td></td>
<td>Middle-aged or elderly males</td>
<td>Leucocytosis &lt;15,000/mm³</td>
<td></td>
</tr>
<tr>
<td>Legionella pneumophila</td>
<td>Onset with flu-like symptoms and chest pain, confusion, diarrhoea and abdominal pain</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mycoplasma pneumoniae</td>
<td>More gradual onset with general symptoms of malaise, body pains and headache, with symptoms of upper respiratory tract infection</td>
<td>Young adults and children</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Chlamydia psittaci</td>
<td></td>
<td>Adults</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coxiella burnetti</td>
<td></td>
<td>Adults</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Influenza viruses</td>
<td></td>
<td>All ages</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Adenovirus</td>
<td></td>
<td>Young adults and children</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Respiratory syncytial virus</td>
<td></td>
<td>Children 3-5 years old</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Measles</td>
<td>Rash, etc.</td>
<td>Children</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Varicella</td>
<td>Rash, etc.</td>
<td>Adults</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table H.2 Factors predisposing to increased risk of venous thromboembolism

<table>
<thead>
<tr>
<th>Disease process</th>
<th>General</th>
</tr>
</thead>
<tbody>
<tr>
<td>Surgery – especially pelvic</td>
<td>Increasing age</td>
</tr>
<tr>
<td>Myocardial infarction</td>
<td>Previous thromboembolism</td>
</tr>
<tr>
<td>Congestive cardiac failure</td>
<td>Varicose veins</td>
</tr>
<tr>
<td>Trauma</td>
<td>Obesity</td>
</tr>
<tr>
<td>Stroke</td>
<td>Pregnancy</td>
</tr>
<tr>
<td>Malignant disease – especially pancreatic</td>
<td>Oral contraceptive use</td>
</tr>
<tr>
<td>Lower limb fracture – especially the hip</td>
<td>Immobility</td>
</tr>
<tr>
<td></td>
<td>• Bed rest</td>
</tr>
<tr>
<td></td>
<td>• Air and bus travel</td>
</tr>
</tbody>
</table>

with pneumococcal pneumonia usually does not appear until several days after the beginning of the illness. If the sputum is (or becomes) frankly purulent, the possibility of a suppurative pneumonia or lung abscess must be considered. Haemoptysis associated with lung abscess may be secondary to bronchial obstruction (e.g. by carcinoma), especially in middle-aged or older smokers, but more rarely by an adenoma in younger adults, or a foreign body in children.

Pulmonary infarction

This diagnosis should be thought of if haemoptysis is preceded or accompanied by pleuritic pain of sudden onset, and possibly by slight fever usually with dyspnoea, and can precipitate atrial fibrillation, or if the patient has present evidence of deep leg vein thrombosis providing an obvious origin for a pulmonary embolus. However, in many patients, signs of pelvic or leg vein thrombosis do not become clinically manifest until days or weeks after an episode of pulmonary embolism, and sometimes never at all. Disorders known to increase the risk of venous thromboembolism – which, if present, should raise the diagnostic suspicion of pulmonary infarction as the cause of haemoptysis – are listed in Table H.2.

Bronchiectasis

Haemoptysis in a patient with a long history of cough and persistently or intermittently purulent sputum, possibly with episodes of increased volume and purulence associated with fever, suggests bronchiectasis (see Fig. D.25). Localized crackles may be found persistently over the affected part of the lung, and clubbing of the fingers may be present, particularly in those with long-standing persistently purulent infection. Bleeding probably arises from the large pulmonary–systemic arterial anastomoses that develop in long-standing cases; it may be the principal symptom and may be severe, even with bronchiectasis of limited extent. CT scanning of the thorax may help to establish the diagnosis.

Chronic bronchitis

Patients with chronic bronchitis not uncommonly cough up blood-streaked sputum, especially during an exacerbation of their condition; expectoration of pure blood is much less common. In either case, some explanation for the haemoptysis, other than chronic bronchitis, should always be considered. Those with chronic bronchitis are nearly always cigarette smokers,
and they are also candidates for carcinoma of the lung. (For a discussion of the recognition of chronic bronchitis, see SPUTUM p. 636.)

**Aspergilloma**
The fungus *Aspergillus fumigatus* particularly colonizes not only open healed tuberculous cavities but also any previously damaged lung tissue, such as occurs in bronchiectasis, sarcoidosis and pneumoconiosis, diffuse fibrosis and localized fibrosis of the lung (as associated with ankylosing spondylitis). The mycelia grow into a ball that almost fills the cavity but leaves a crescent of air above the opacity. This results in the characteristic radiological appearances.

Aspergillomas are usually discovered on a routine chest X-ray, but recurrent haemoptysis is a characteristic feature and may herald a massive pulmonary haemorrhage.

A sputum examination for aspergilli is unhelpful because the organisms may be present in healthy persons and are, in any case, not often identified in the sputum of patients with aspergilloma. There is a useful precipitin test that is almost always strongly positive, whereas the aspergillin skin test is positive in only about 20 per cent of cases.

**Foreign bodies in the tracheobronchial tree**
These can give rise to haemoptysis in two ways. Soon after lodgement, a hard foreign body with sharp edges may lacerate the mucosa, cause local ulceration and lead to bleeding, which is usually slight, with no more than blood-streaked sputum. Later, infection beyond a foreign body obstructing a bronchus can cause pneumonia, abscess formation and, if neglected, eventually bronchiectasis – all of which are possible causes of haemoptysis. If it is not radio-opaque, or if it is lodged centrally where it may be hidden in the mediastinal shadow, a foreign body will be radiologically inapparent and provide no radiological evidence of its presence until the secondary changes arising from bronchial obstruction appear. A non-occluding foreign body of metal, bone or plastic material often produces no immediate irritative symptoms, and haemoptysis may be the first symptom to draw attention to its presence. The diagnosis will be made or confirmed at bronchoscopy.

**Hydatid cyst**
Infection with *Echinococcus granulosus* is particularly prevalent in the Middle East, Mediterranean coastal countries, South America, South Africa and Australia. Haemoptysis is the most common single symptom of an intact hydatid cyst and is possibly due to the size of the cyst causing congestion and blood vessel erosion; indeed, it often precedes cyst rupture. When a cyst has ruptured, infection often complicates the picture, with all the possible consequences of pulmonary suppuration, including severe haemoptysis. The diagnosis will be suggested by the chest X-ray, which will show one or more rounded shadows. The rupture of cysts and added infection will, of course, affect the appearances. Complement fixation and Casoni skin tests may be helpful in diagnosis.

**Paragonimiasis**
Haemoptysis is a leading feature of infection with the lung fluke, *Paragonimus westermani*. This occurs endemically in Japan, China, Korea and Taiwan; it also occurs in parts of Central and West Africa. In addition to haemoptysis, cough and chest pain are apparent. The chest X-ray shows characteristic air-containing cysts, usually 1–2 mm in diameter with a thickened base, scattered throughout the lungs. These cysts may be mistaken for tuberculous cavities, particularly if the changes are limited to the subclavicular zones as frequently happens. The diagnosis finally depends upon the finding of ova in sputum or stool.

**Other parasitic infestations**
Haemoptysis may occur either as parasites pass through the lungs or when they finally settle in the lungs. Most of these parasites have fairly well-defined geographical distributions and are only likely to affect those patients who are (or have been) resident in endemic areas. Among parasites that may cause haemoptysis by their passage through the lungs are included:

- *Ascaris lumbricoides*, which has a worldwide distribution
- *Schistosoma*, of which various species have different, mainly tropical, distributions. These cause haemoptysis during the passage of larvae through the lungs, although the principal pulmonary manifestation – obliterative arteriolitis with granuloma formation – arises later and is not especially associated with haemoptysis
- *Dirofilaria immitis*, the heartworm of dogs, which may cause haemoptysis and radiologically detectable lesions in the lungs of humans

**Pulmonary haemosiderosis**
Pulmonary haemosiderosis is a condition that is characterized by episodic bleeding into the lungs, and this results in the deposition of haemosiderin in intra-alveolar macrophages and in interstitial histiocytes, together with a variable degree of alveolar wall fibrosis.
in the more chronic forms of the disease. The disorder presents in different ways at different ages.

In childhood haemosiderosis, acute episodes of pulmonary haemorrhage with fever, cough, haemoptysis and breathlessness occur, and the child is found to be anaemic. The individual episodes are often self-limiting and clear rapidly, but they can be life-threatening. The sputum characteristically contains iron-laden macrophages between the episodes of haemorrhage. Physical signs in the chest are unimpressive; crackles are sometimes heard. The chest X-ray shows large bilateral confluent lesions that appear and disappear rapidly. Antibasement membrane (ABM) antibodies are not found.

In adult idiopathic haemosiderosis, there is a gradually increasing breathlessness accompanied by small ‘fleck’ haemoptysis, which occurs repeatedly over a prolonged period. The chest X-ray shows widespread fine stippling, often of pinhead size, and most densely distributed at the bases. Areas of acute confluent shadowing are rarely seen. The appearance may remain unaltered over many years. A physical examination of the chest usually reveals little in the way of abnormal signs unless alveolar wall fibrosis is marked, when finger clubbing may develop. The diagnosis is made by finding haemosiderin-laden macrophages in the sputum between episodes of haemoptysis. If there is no sputum for examination, macrophages can be obtained by bronchoalveolar lavage using the fiberoptic bronchoscope. Large numbers of these iron-laden cells are obtained, sometimes rendering the lavage fluid almost black. Some patients have airflow obstruction as well as a restrictive defect; the explanation for this is not known, but the finding does not negate the diagnosis.

Diffuse lymphangioleiomyomatosis

This exceedingly rare condition is diagnosed by histological examination of a lung biopsy from patients who present with a diffuse lung disorder, haemoptysis and slowly increasing breathlessness. Obstructive distortion of the pulmonary veins by hypertrophic muscle leads to capillary haemorrhage and resulting siderosis. Other clinical features are recurrent pneumothoraces and chylothorax. The aetiology has not been established, but the condition is probably a hamartomatous malformation of lymphatic and perilymphatic tissue.

Systemic vasculitides

In the rare cases of polyarteritis nodosa in which the pulmonary arteries are involved with consequent small infarctions of the lung, haemoptysis may occur. Suspicion of the diagnosis depends on the other systemic features of polyarteritis nodosa. Confirmation of the diagnosis may be by tissue biopsy or the demonstration of multiple intra-parenchymal aneurysms on visceral angiography.

In the Churg–Strauss syndrome (eosinophilic granulomatosis), there is usually a long history of asthma with eosinophilia, and eventually the patient develops other manifestations of generalized vasculitis. Haemoptysis, although not a particular feature, may occur; the chest X-ray may be normal or show transient or ‘fixed infiltrations’ or nodules.

Haemoptysis is more likely to be a clinical symptom in Wegener’s granulomatosis – now more correctly referred to as granulomatosis with polyangiitis. This condition is characterized by necrotizing granulomas in the respiratory tract, including the nose and sinuses. There is also widespread vasculitis that usually affects the kidney. The chest X-ray shows multiple, usually bilateral nodular shadows that may vary in size from time to time and may cavitate; when they do so, the cavities often have thick irregular walls. Diagnosis is made by biopsy of a lesion in the nose, sinuses or lung. Goodpasture’s syndrome occurs most frequently (but not exclusively) in young male cigarette smokers (male:female ratio 6:1). The pulmonary lesions usually develop first; there is episodic haemoptysis that is sometimes severe enough to be life-threatening, and the chest X-ray shows confluent or widespread fine shadows related to the severity of the bleeding. Within about a year, renal impairment develops, with fairly rapid progression to renal failure and death.

The feature that distinguishes this form of pulmonary haemosiderosis from the others is the presence in the serum of ABM antibodies. These antibodies are believed to cross-react with the basement membranes of both the lungs and the kidneys. Renal and lung biopsies demonstrate a pattern of linear fluorescence, both in the glomeruli and along the capillaries in the alveolar walls. Typically, the disorder affects cigarette smokers who present with cough, dyspnoea and haemoptysis that may be streaky and intermittent, or sometimes severe enough to be life-threatening. The lungs are usually spared in non-smokers.

Systemic lupus erythematosus (SLE) may be associated with recurrent haemoptysis. The chest X-ray usually shows widespread confluent patchy shadows, and the patient has other clinical features of SLE; immunological tests confirm this diagnosis. The lesion in the lung is believed to be due to a pulmonary vasculitis. In contrast to some other types of haemosiderosis, resolution with corticosteroids is often dramatic.
Arteriovenous fistula of the lung
This usually causes symptoms and signs arising from shunting of mixed venous blood directly from pulmonary artery into pulmonary vein – effectively a ‘right-to-left’ shunt. Haemoptysis occurs in only a minority of cases. About half of these are associated with hereditary haemorrhagic telangiectasia, of which epistaxis is a frequent symptom.

Trauma
Haemoptysis occurs not only when the lung is directly penetrated, or lacerated by fractured ribs, but also in non-penetrating injuries. These may be associated with contusion of the lung, even without rib fractures. Exposure to blast from explosions may cause haemorrhagic consolidation of the lung (blast injury) with haemoptysis.

Mitral stenosis
The heart should be carefully auscultated for signs of mitral stenosis in all cases of haemoptysis. Rheumatic heart disease is less often encountered in Western countries. The cause of the haemoptysis may be the high pulmonary venous pressure or pulmonary embolism and infarction, particularly in patients with atrial fibrillation and/or cardiac failure. Pulmonary embolism leading to infarcts, especially in patients with atrial fibrillation, is a frequent cause. Less severe bleeding leading to bloodstained sputum may be associated with the high pulmonary venous pressure.

Left ventricular failure
The thin, frothy sputum produced in pulmonary oedema is frequently tinged pink with blood. Diagnosis depends upon the recognition of the underlying cardiac disease.

Aortic aneurysm
An aortic aneurysm may erode into a bronchus, leading to rapidly fatal haemorrhage with massive haemoptysis. Most aneurysms of the thoracic aorta seen now arise as a result of traumatic dissection following a road traffic accident. The classic syphilitic aortic aneurysm has become a rarity.

Bleeding diathesis
In disease associated with disturbances of haemostasis and clotting, such as thrombocytopenia, Henoch–Schönlein purpura, scurvy, leukaemias and aplastic anaemia, haemoptysis may occur, but as a minor feature of a generalized bleeding tendency in which epistaxis and bleeding from the gums are more prominent. These may be accompanied by bleeding from the alimentary or urinary tract, or by purpura, leading to appropriate haematological investigations. Haemoptysis is virtually never the sole clinically evident presenting feature of these diseases.

Factitious ‘haemoptysis’
When a patient presents with a history of haemoptysis as the sole symptom, and investigation has shown no evidence of its source or of any organic disease, it must be remembered that single episodes of unexplained haemoptysis are not uncommon. Moreover, the patient may have come for investigation because they are seeking reassurance; for instance, they may have a friend who has been found to be suffering from tuberculosis or bronchial carcinoma with haemoptysis as an initial symptom, and have become alarmed lest the streaks of blood they notice after cleaning their teeth are the first sign of the same thing in themselves. Very occasionally, a patient returns with recurrent complaints of haemoptysis, the blood being deliberately produced by various forms of trauma in the mouth or pharynx.

Hallucinations
Andrew Hodgkiss
Hallucinations may most simply be defined as percepts without objects. They need to be distinguished from illusions, which are misperceived objects.

It has been known for well over a century that certain types of hallucination are found in normal mental health. For example, hearing one’s name called in the street or hearing a phone ring is not unusual. States of expectancy, exhaustion, sensory deprivation or sleep deprivation will certainly provoke hallucinations in people with good mental health. In such instances, a degree of insight into the hallucinatory nature of the perception is generally maintained, and some authorities call such phenomena ‘pseudohallucinations’. Bereavement is another state in which hallucinations commonly occur, typically auditory or visual hallucinations of the deceased. Brief hallucinations on falling asleep (hypnagogic) or upon waking from sleep (hypnopompic) are also reported in the general population, and this can also be prominent in narcoleptic patients.

However, the sensory modality, duration and other detailed features of hallucinations can be of great diagnostic value. For example, if a general hospital inpatient describes visual hallucinations, they are best initially considered delirious until proven otherwise. Intoxication or withdrawal from drugs or alcohol must be immediately considered. Delirium tremens, an acute confusional state
complicating the early days of withdrawal from alcohol, typically includes visual hallucinations of insects or rodents scuttling along floors and walls. Prescribed medication with atropine-like actions is a potent cause of visual hallucinosis and disorientation in the elderly. If the visual hallucination recurs sporadically over several weeks but is always the same in form and content, the possibility of an epileptic focus or space-occupying lesion is raised. Sustained Lilliputian visual hallucinations are almost pathognomonic of the rare Charles Bonnet syndrome.

About 20 per cent of patients with schizophrenia report visual hallucinations, and these can be quite complex tableaux connected to delusional beliefs. Visual hallucinations in psychotic depression are rare and tend to be mood-congruent pictures of morbid scenes such as coffins and corpses. Finally, the classical visual hallucinations of LSD intoxication (and subsequent flashbacks) deserve mention. These tend to involve shifting geometric patterns or surreally altered figures (e.g. a man with the head of a tortoise).

Auditory hallucinations can also be of great diagnostic value. Patients who hear their own thoughts spoken out loud in external space, or their thoughts echoed by voices, or their behaviour discussed by others in the third person as a running commentary, are most likely to suffer from schizophrenia. In the acute, early phase of the disorder, this can be an extremely distressing experience, while over a period of years many patients gain a degree of acceptance or even mastery experience, while over a period of years many patients gain a degree of acceptance or even mastery of the problem. It should be added that auditory hallucinations of this form are also found in 20 per cent of manic patients. The auditory hallucinations of patients with psychotic depression or mania tend to be briefer, mood-congruent phenomena in the second person (for example ‘why don’t you just kill yourself?’, or ‘you are the Messiah’). Auditory hallucinations do occur in delirium, and musical hallucinations in particular should prompt a serious investigation for an organic aetiology.

Olfactory hallucinations, typically of unpleasant smells, occur commonly in schizophrenia but also as an aura to temporal lobe epilepsy. Tactile hallucinations, particularly formination (the feeling of ants crawling under the skin), point to cocaine misuse or alcohol withdrawal. Bizarre somatic hallucinations, such as the feeling of being impregnated, do occur in schizophrenia.

The common causes of hallucinations are summarized in Box H.4.

### Box H.4 Causes of hallucinations

- Sleep and sensory deprivation
- Extreme fatigue
- Bereavement
- Narcolepsy
- Delirium
- Alcoholic hallucinosis
- Drugs
  - Amphetamine
  - Cocaine, LSD, mescaline, magic mushrooms, dimethyltryptamine, anticholinergics, bromocriptine, inhaled solvents
- Schizophrenia
- Mania
- Psychotic depression
- Post-concussional states
- Temporal lobe epilepsy
- Intracranial space-occupying lesions

### HEAD, RETRACTION OF

Mark Kinirons

The common causes of head retraction (and stiffness) are listed in Box H.5. At its most dramatic (as in meningitis), it is possible to lift the whole body by lifting the head up. Stiffness due to disease of the cervical spine and para-spinal tissues is dealt with elsewhere (see NECK PAIN AND/OR STIFFNESS, p. 452). The term meningism is applied to the headache, photophobia and stiff neck that occur, as a rule, in children, in the course of general infections such as tonsillitis, pneumonia and pyelitis. The pressure of the spinal fluid is raised, but its contents are normal.

Meningitis causes resistance to forward flexion of the neck, but this may be absent in very mild cases and also in fulminating infections. Actual retraction of the head is best seen in tuberculous meningitis and in meningococcal meningitis. Inflammation of the leptomeninges is caused by many organisms – bacteria, viruses, spirochaetes and yeasts – and a low-grade ‘meningitis’ can occur when the meninges are invaded by secondary carcinomatosis and sarcoidosis.

Features common to most cases of meningitis are headache, photophobia, vomiting, giddiness and fever. There may be a rigor, or a convulsion, at the onset of the more virulent types, especially in children. There is stiffness of the neck, spinal muscles and hamstrings. Thus, forward flexion of the neck is resisted, and it may evoke flexion of the hips and knees (Brudzinski’s sign). There is resistance to extension of the knee on the flexed thigh (Kernig’s sign), because this movement pulls on the roots of the cauda equina.
Meningococcal meningitis

Meningism

Meningitis

– Bacterial

– Viral

– Spirochaetal

– Fungal

– Carcinomatous

– Sarcoid

– Subarachnoid haemorrhage

– Pressure cones

– Asphyxia

– Intermittent retraction

– Spasmode tocricollis

– Torsion spasm

– Tetanus

– Rabies

– Streptococcal poisoning

– Spinal and paraspinal disease

There may or may not be evidence of focal damage to the brain and cranial nerves. The latter are involved as they traverse the subarachnoid space, and the brain itself can be damaged by spread of the infection along the meningeal sheaths that cover the vessels as they penetrate the surface of the brain. Moreover, thrombosis both of arteries and of veins can occur, with infarction, oedema or brain abscess as a result.

Infection can gain access to the meninges by several routes. In most cases it is blood-borne, and in an important minority it spreads from local infections in the ear, accessory nasal sinuses, face and scalp. The existence of this second group emphasizes the need to seek for evidence in the history and on physical examination as to the possibility of local infection in every case. Points to be looked for are the presence of otitis, mastoiditis or sinusitis. Moreover, a history of head injury, whether recent or remote, may mean that there is a fracture and a dural tear leading into an air sinus, thereby providing a path for the entry of microorganisms. In such cases, meningitis is apt to be associated with an abscess, whether extradural, intradural or intracerebral, along the track of entry.

Sepsis in the face or scalp, such as furunculosis, erysipelas, infected scalp wounds or herpes, can lead to meningitis in debilitated persons. Infection may also enter via a meningocele, or a congenital dermal sinus at the base of the spine, and these must be looked for in unexplained meningitis in infants. Yet another manner of infection is following lumbar puncture or spinal anaesthesia, fortunately rare; in such cases low-grade infection is usual (e.g. by Bacillus pyocyaneus).

Meningococcal meningitis (spotted fever, cerebrospinal fever) usually occurs in epidemics that are initiated by droplet infection from healthy carriers. A bacteraemia precedes the meningitis by hours, days or even weeks. Occasionally, there is a chronic meningococcal septicaemia with fever, purpura and transient pain, and swelling in the joints, and a proportion of such cases will end up with meningitis. In another small group, the patient is overwhelmed by a fulminating septicemia within a few hours of the onset; some pass rapidly into coma, without a significant fall in blood pressure, while others remain clear in mind but suffer a drastic fall in blood pressure due to circulatory collapse, and these cases usually present a diffuse purpuric rash on the skin (the Waterhouse-Friderichsen syndrome, which is also seen in other severe infections) (see Fig. P.47, p. 562). In the usual type of meningococcal meningitis, however, there is fever, meningeal irritation, severe headache and sometimes a purpuric or macular rash. Convulsions may occur at the onset. Transient cranial nerve palsies and papilloedema may be found, and delirium is common. Tendon reflexes are reduced, and extensor plantar responses are common in the more severe cases. The spinal fluid in meningitis usually contains a polymorph pleocytosis, with a rise of protein and a fall of glucose. Both intracellular and extra-cellular diplococci are found.

In some cases of meningococcal meningitis, the exudate is largely confined to the base of the brain, thereby leading to an obstructive hydrocephalus. There is mild fever, vomiting, papilloedema and head retraction. In infants (the usual victims), the head enlarges and there is a slow downward course with emaciation, vomiting and increasing stupor. During the first few days of the illness, the changes in the spinal fluid are the same as in the ordinary type of meningococcal meningitis, but thereafter the meningococci disappear and there is merely a lymphocytic pleocytosis, a rise of protein, and rather a low sugar content. In such cases, ventricular tap may produce the diplococcus.

Further sequels of meningococcal meningitis are the subdural hygroma referred to above, cranial nerve palsies, and disabilities arising from the formation of scar tissue around the spinal cord. These include a lower motor neurone paralysis of muscles in the limbs, and occasionally an incomplete transverse lesion of the cord with paraplegia, sensory impairment and sphincter disturbances.

Other forms of pyogenic meningitis are sporadic rather than epidemic in incidence, do not as a rule produce a rash, and are usually derived from a more or less obvious source of infection. Thus, pneumococcal cases commonly arise from infection in the ears or sinuses, or from pneumococcal pneumonia. Streptococcal meningitis is rarer than the pneumococcal form but occurs in similar circumstances; in a proportion of cases, however, there is a cerebral abscess in addition to the meningitis, and this must always be looked for,
Meningitis with a predominantly lymphocytic response in the cerebrospinal fluid (CSF) occurs in infection by tuberculosis, viruses, yeasts and spirochaetes. Of these, tuberculosis is the most important. The meningitis is secondary to tuberculosis elsewhere, although the source may not be clinically apparent. It rarely occurs before the age of 6 months, and it is most common in children and young adults. The onset is commonly insidious, with malaise and occasional headaches that may precede the signs of meningitis for days, weeks or even months. The meningeal phase includes headaches, signs of meningeal irritation and retraction of the head. Epileptic attacks – whether focal or general – and sudden hemiplegia or monoplegia, aphasia or cranial nerve palsies, mental changes, and papilloedema with visual loss, are common; hydrocephalus tends to increase, leading in untreated cases to stupor, incontinence, a rise in pulse and medullary failure. Choroidal tubercles may be found on examination of the retina, and the Mantoux test is positive in the majority of cases. The spinal fluid is under increased pressure and may be either clear or opalescent; a fibrin clot forms on standing for some hours. There is an excess of lymphocytes, and there may be a few polymorphs. The protein is raised, and the sugar content falls at an early stage. The definitive test is the demonstration of the organism in the fluid by direct smear with Ziehl–Neelsen staining; if the evidence in favour of the disease is good, treatment should not be withheld until the organism is found. In the early stages of the disease, the conditions that may cause difficulty in diagnosis are acute lymphocytic choriomeningitis and other virus infections of the meninges in which, however, the sugar and chloride content of the CSF are normal, and the clinical course is quite different, with rapid recovery in most cases. In acute syphilitic meningitis, the VDRL reaction is positive, while in meningitis associated with aural and sinus infections, both lymphocytes and polymorphs may be present, with negative culture and a normal sugar content in some cases (aseptic meningitis).

Meningitis can complicate Weil's disease (spirochaetosis icterohaemorrhagica), but an acute and predominantly lymphocytic meningitis can also occur without jaundice, renal damage or haemorrhagic symptoms – a syndrome complex that is strongly suggestive of this spirochaete. Weil's disease occurs in persons who have been in contact with rats (e.g. canal-bathers, sewage workers, etc.), and the CSF is sterile if ordinary culture media are used. Sugar and chloride contents are normal. A benign meningitis can also be caused by Leptospirosis canicola, which is carried by dogs. There may be conjunctival suffusion, and a rash that may resemble erythema nodosum. The CSF contains an excess of lymphocytes, while the sugar content is normal, and the fluid is sterile in ordinary culture media. Diagnosis is confirmed, as in Weil's disease, by guinea-pig inoculation and by the detection of antibodies in the blood.

Lymphocytic meningitis may also occur with tick-borne relapsing fever, either during the first attack of fever, or more often in subsequent bouts. There is severe headache, neck stiffness and slight papilloedema. Cranial nerve palsies, notably of the VIth nerve, are not uncommon. There is increase of protein and lymphocytes in the CSF, and the organism can be identified by dark-ground illumination, or by inoculation of the CSF into a suitable animal. A similar syndrome occurs with Lyme disease due to Borrelia burgdorferi with the typical addition of a skin rash.

A well-marked lymphocytic meningitis can be caused by the viruses of acute choriomeningitis, mumps and glandular fever, whereas the meningeal reaction of poliomyelitis, zoster and arthropod-borne encephalitis is usually less obtrusive. A specific virus is responsible for acute lymphocytic aboriomeningitis, a benign disease characterized by a prodromal period of malaise, headaches, muscle pains, pyrexia and upper respiratory catarrh; this is followed after a week or two by severe headache, photophobia, neck stiffness and a positive Kernig's sign. In a minority of cases, transverse myelitis, facial palsy or temporary mental and emotional changes may occur. The protein level is raised in the CSF, and there may be from 50 to 3000 cells/ml of CSF, of which at least 95 per cent are usually lymphocytes. Sugar is normal, and the virus can sometimes be isolated from the CSF. Mumps meningitis usually starts on the fifth to tenth day of the illness, but meningeal symptoms may precede the parotitis, or they may occur with orchitis but without parotitis. The meningitis may be accompanied by encephalitis with disturbances of consciousness and, rarely, focal cerebral and cerebellar signs. Cranial nerve palsies and myelitis have been described. Sudden permanent deafness in one or both ears, and with or without vertigo and vomiting, may occur in...
mumps without evidence of meningoencephalitis. In glandular fever, there may be a well-marked lymphocytic meningitis of sudden onset, with enlargement of the glands, an increase in mononuclear lymphocytes in the blood, and an increasing titre in the Paul–Bunnell test. Acute polyneuritis may complicate the disease, or it may occur in glandular fever without meningitis. Japanese B encephalitis occurs in Japan and China. The virus is spread by mosquitoes. There are meningeal signs, drowsiness, stupor, signs of diffuse cerebral involvement, convulsions and tremors. There is a lymphocytosis in the CSF, while the sugar remains normal. The mortality in this disease can be over 50 per cent.

Infection by yeasts has been uncommon in the past, but it appears to be on the increase since the advent of antibiotics. Cryptococcosis (Torula histolytica) involves the subcutaneous tissues, the lung and the central nervous system, either alone or in combination, or in series. Subcutaneous granulomas break down to form abscesses and ulcers. Pulmonary lesions may mimic either chronic tuberculosis or carcinoma. The cerebral type usually starts insidiously with headaches, dizziness and stiffness of the neck, but it may commence suddenly. There is little or no fever, but gradually the CSF pressure rises, producing papilloedema, and there may be cranial nerve palsies, hemiparesis or ataxia. Large granulomas may, in fact, cause symptoms of a cerebral tumour. The patient eventually sinks into coma. There is a marked mononuclear pleocytosis in the CSF, and the protein is raised. The glucose content is reduced, and cryptococci, which are readily mistaken for erythrocytes or lymphocytes, can be found in small numbers in the CSF.

Sarcoidosis, which causes uveo-parotid polyneuritis, can also give rise to a low-grade meningitis with headaches, slight stiffness of the neck, and a rise in lymphocytes and protein in the CSF. It can pass on to cause an obstructive hydrocephalus, with papilloedema and optic atrophy. Cranial nerve palsies and diabetes insipidus have been described. The diagnosis can only be inferred by the presence of typical lesions in other areas (e.g. skin, liver, lungs and eyes).

Subarachnoid haemorrhage is usually due to rupture of a saccular aneurysm, or of an atheromatous aneurysm. Less common causes are hypertension, angiomatic malformations, mycotic and syphilitic aneurysms, and purpura. An abrupt onset, early loss of consciousness in most cases, and the presence of blood in the CSF distinguish the average case from meningitis. When the leak is slow, however, the severe headache, stiffness of the neck, slight pyrexia, ocular palsies and positive Kernig’s sign may simulate meningitis, and it is only the blood in the CSF that clinches the diagnosis. If the lumbar puncture is delayed for a day or two, the CSF may be found to be yellowish in colour (xanthochromia), and not bloodstained. Therefore, xanthochromia should be sought if subarachnoid haemorrhage is suspected. Rarely, pain starts in the lumbar region and gradually spreads down the back of the legs and up to the neck. Another unusual form presents as sudden coma, and there may, in such a case, be a history of former attacks of unexplained coma with neck stiffness.

Pressure cones at the tentorial hiatus and at the foramen magnum can cause stiffness of the neck. They occur as a result of space-occupying lesions and, occasionally, from cerebral oedema. The local rise of pressure from an expanding mass in the head, or from hydrocephalus, dislocates and displaces brain substance. Thus, a mass in the middle fossa (e.g. tumour or extradural haematoma) can dislocate part of the temporal lobe into the posterior fossa, with the result that the midbrain and the displaced tissue are tightly wedged in the dural ring (Fig. H.7).
HEADACHE

David Werring & Mark Kinirons

Headache (cephalgia) is one of the most frequent reasons for visiting a doctor. It is not known why headache is so common, but the face, scalp, nasal passages, eye and ear contain many pain receptors, and humans are concerned about pain in the head due to the possibility of a serious cause (e.g. a brain tumour). In the UK, about 80 per cent of the population will experience a headache in a given year. About 1–2 per cent of the population will consult their general practitioner, and about 0.3 per cent will be referred on to a hospital specialist, usually a neurologist. The vast majority of headaches seen in primary care (over 95 per cent) are not due to a serious intracranial cause. There are many possible causes of headache (Box H.6), and the first priority is to make sure that serious or treatable intracranial pathology are not missed. In general, the length of the history is a useful guide: headache with a short history requires prompt diagnosis and possibly urgent investigations.

This may obstruct the aqueduct, thus aggravating the situation by causing internal hydrocephalus. In posterior fossa tumours, the reverse is seen: the oedematous brainstem and cerebellar tissue is displaced upwards through the tentorial notch. Downward herniation of the medulla and cerebellar tonsils through the foramen also occurs, and this will also give rise to rigidity of the neck.

Pressure cones can arise as the result of intracranial space-occupying lesions: tumour, abscess, haematoma, internal hydrocephalus and occasionally cerebral oedema due to vascular lesions. All of these conditions may therefore cause stiffness of the neck. It is important to recognize the presence of a pressure cone, because the removal of even a small quantity of CSF by lumbar puncture may cause collapse and death.

Asphyxia can cause retraction of the head, or stiffness of the neck. The more striking examples are usually seen in children with bronchopneumonia, bronchiolitis or foreign body in the larynx. Asphyxia has also sometimes been noted in retropharyngeal abscess. Even in adults with severe bronchopneumonia, there may be stiffness of the neck, although retraction is rare. That asphyxia without cerebral oedema can cause retraction of the head is well illustrated by the retraction seen during the administration of pure nitrous oxide (e.g. for dental extraction), but this may not be the whole explanation in the diseases mentioned.

Other features indicating the possibility of serious intracranial pathology are: sudden onset of pain (possible intracranial haemorrhage); fever or neck stiffness (possible intracranial infection); focal neurological symptoms or signs (possible mass lesion); and temporal artery tenderness (possible giant-cell arteritis).

Although the clinician must be alert to serious causes of headache, it should be remembered that in primary care fewer than 5 per cent of headaches are due to serious intracranial pathology. Migraine and tension-type headache are by far the most common causes seen in clinical practice. A detailed history should allow an accurate diagnosis of these types.

Box H.6 Causes of headache

- Migraine
  - Migraine with aura
  - Migraine without aura
  - Special forms (ophthalmoplegic migraine, retinal migraine, acephalagic migraine)
- Tension-type
- Trigeminal autonomic cephalgias
  - Cluster headache
  - SUNCT (short-lasting unilateral neuralgiform headache with conjunctival injection and tearing)
  - Paroxysmal hemicrania
- Vascular disorders
  - Acute ischaemic stroke
  - Acute haemorrhagic stroke (subarachnoid or intracranial haemorrhage)
  - Reversible cerebral vasoconstriction syndrome (RCVS)
  - Temporal arteritis
  - Hypertension
  - Venous sinus thrombosis
- Non-vascular intracranial pathology
  - Abnormal cerebrospinal fluid pressure (high or low)
  - Infection (meningitis, meningoencephalitis, abscess)
  - Tumours
  - Systemic infection (viral, bacterial, others) or metabolic disturbance
- Head and face pain arising from other structures
  - Cranial bones, neck, eyes, ears, nose, sinuses, teeth, jaws, temporomandibular joints
- Craniofacial neuralgias
  - Trigeminal neuralgia
  - Glossopharyngeal neuralgia
- Specific headache syndromes
  - Idiopathic stabbing headache
  - Coital cephalgia
  - Cold stimulus headache (ice-cream)
  - Benign exertional and cough headache
- Trauma
- Substance use
- Substance withdrawal
of headache, but this often requires considerable patience in allowing the history to unfold. It should be evident that, because headache can be due to serious pathology, symptomatic treatment should never precede a careful history, adequate physical examination and consideration of further investigation. As in all types of pain, the quality, location, severity, time course and exacerbating and relieving factors should be determined when assessing headache. The head should be examined for signs of temporal arteritis (pulseless, thickened, tender temporal arteries). The blood pressure should be measured and the optic fundi examined for papilloedema, which usually indicates raised intracranial pressure (papilloedema can also be caused by local pathology at the optic nerve head; see OPTIC FUNDUS, ABNORMALITIES IN, p. 476). Raised intracranial pressure may be due to a structural lesion, or to obstruction of normal cerebrospinal fluid (CSF) circulation, as in idiopathic intracranial hypertension or cerebral venous sinus thrombosis. A neurological screening examination should be performed, to include pupils (including looking for Horner’s syndrome), visual fields, extraocular movements, drift of outstretched hands, coordination testing (finger–nose test), reflexes and plantar responses, and heel–toe walking; recent-onset headache with focal neurological signs is an indication for urgent brain imaging.

The differential diagnosis of headache is shifted at different ages; for example, cranial arteritis is almost exclusively seen in the elderly (aged 60 years and above), whereas in children posterior fossa tumours must be considered with any refractory recent-onset headache. In this discussion, the common causes of headache (migraine and tension-type) will be considered first and in detail, before dealing with headache due to vascular causes (intracranial and extracranial), non-vascular intracranial pathology including tumour, infection (systemic or intracranial), and finally some rare but distinctive headache syndromes and headaches due to systemic disease. Pain which is mainly limited to the face (craniofacial neuralgias; head or face pain due to disorders of teeth, sinuses, ear, nose and other structures) is considered elsewhere (see FACE, PAIN IN, p. 182).

MIGRAINE
Pathophysiology
A useful pathophysiological model of migraine is that of an inherited disorder causing increased sensitivity to afferent stimuli; during attacks, stimuli including light, sound and movement exacerbate the headache. Migraine attacks are thought to result from increased neuronal activity in midline brainstem structures, including the periaqueductal grey matter, the dorsal raphe nuclei, reticular formation and locus coeruleus. Functional imaging studies show increased cerebral blood flow in these areas during an attack. These brainstem nuclei, together with the trigeminal nuclei, can influence extracranial blood flow by reflex connections with the parasympathetic part of the facial nerve. This link between neural and vascular systems is termed the trigeminovascular system. The same brainstem nuclei can also modulate cortical activity and blood flow, as well as central pain control mechanisms. The migraine ‘aura’ is thought to involve a wave of neuronal depression spreading across the cortex, associated with reduced cerebral blood flow (oligemia) and neurological symptoms corresponding to the brain region affected. Thus, the visual aura may propagate across the visual field as the wave of cortical spreading depression (CSD) moves across the occipital cortex. The pain is thought to result from neurogenic inflammation resulting from the release of neuropeptides (by trigeminal nerve endings) and other pain-inducing compounds including histamine, serotonin, prostaglandins and nitric oxide (from plasma, platelets and mast cells). These agents induce cranial vasodilatation and extravasation of plasma proteins, as well as the sensitization of trigeminal nociceptive nerve endings. Most current treatments for migraine act on the serotonergic system.

Clinical features
Migraine causes severe episodic headaches that last from several hours to several days, and that are absent more than they are present. The periodicity is important: attacks typically occur about once per month, and if headaches occur more than twice a week, episodic migraine (on its own) is unlikely to be the diagnosis. In migraine with aura (see AURA, p. 41) – which affects about one-fifth of patients – the aura evolves over minutes and usually lasts under 30 minutes, almost always under an hour. The aura most often arises from the occipital cortex so is commonly visual, involving positive phenomena such as photopsia (unformed flashing lights) or fortification spectra (tessellated structures resembling zigzag lines). Fortification spectra are so named because they resemble the fortifications of a medieval town. Teichopsia is an equivalent term derived from the Greek word teichos (meaning ‘wall’). Negative phenomena (scotomata) may be reported, sometimes following the teichopsia as it propagates across the visual field. Auras may also affect sensation, movement, cognitive or vestibular functions.
Illusions of body image may be reported (‘Alice in Wonderland’ syndrome). The headache typically follows the aura, but less commonly can come first or coincide with it.

The headache in migraine is usually (but not always) unilateral and throbbing. It is severe and made worse by moving around, loud noises or bright light. The patient typically describes wanting to lie down in a quiet, darkened room; they often feel nauseous and may vomit. In migraine without aura (formerly termed common migraine, affecting about 75 per cent of patients), the headache is similar in character to migraine with aura, but fewer specific accompanying symptoms occur than in classical migraine. There may be gastrointestinal symptoms or heightened sensory perception.

Treatment of migraine is of two types: (i) the symptomatic treatment of acute attacks; and (ii) prophylactic agents, which are usually recommended if more than three attacks occur per month. First-line acute treatments include high-dose aspirin with an antiemetic (e.g. domperidone), and the triptans (e.g. sumatriptan) given by the oral or subcutaneous routes. Prophylactic treatments include propranolol and other beta-blockers, and amitriptyline.

Differential diagnosis

The main differential diagnoses are tension-type headache, cluster headache and other trigeminal autonomic cephalgias, and medication overuse headache. The main feature distinguishing it from tension-type headache is its periodicity. Migraine is generally episodic and rarely more frequent than twice a week, whereas tension-type headache is present on most days. It should be noted, however, that migraine and tension-type headache often co-exist, causing so-called mixed chronic daily headache. Furthermore, it is now appreciated that there is a frequent form of migraine (chronic migraine).

The main difference between migraine and the trigeminal autonomic cephalgias (including cluster headache) is that autonomic features (e.g. nasal congestion, conjunctival injection, ptosis and lacrimation) are more prominent in the latter group. Medication overuse headache is more difficult to diagnose; it often results in the context of migraine with regular use of high-dose analgesics, with subsequent transformation into a mixed migraine and chronic tension-type headache disorder.

TENSION-TYPE HEADACHE

As discussed above, the traditional classification of the most common recurrent headache syndromes into migraine and tension-type headache has been recently questioned, and it has been suggested that they are different expressions of the same pathophysiological process. At present, the distinction between them is useful in that different management strategies are effective for each group. Tension-type headache – also referred to as chronic daily headache – is common, accounting for 70 per cent of referrals to a headache clinic. In comparison to migraine, it has no defining characteristics: the headaches are rather featureless, with no photophobia, phonophobia, nausea or vomiting. In tension-type headache, pain is described in many different ways; it is often diffuse, but it may localize to the vertex, forehead or the neck. It is more often bilateral than unilateral. The classic description is of a ‘tight band around the head’ or ‘like the head being in a vice’. Patients will sometimes report that ‘my head feels as if it is bursting’, or that ‘sharp knives are being driven in’. The headache is worse in the evenings, and with fatigue or stress. Because the headache is chronic, often occurring for years, analgesic misuse is a frequent problem. The most effective drug treatment for recurrent tension-type headache is with tricyclic agents (e.g. amitriptyline). Relaxation exercises may also be helpful.

CLUSTER HEADACHE AND THE TRIGEMINAL AUTONOMIC CEPHALGIAS

Cluster headache

Cluster headache is one of the trigeminal autonomic cephalgias, a group of primary headache disorders linked by their trigeminal distribution, short duration and prominent ipsilateral cranial autonomic features. Cluster headache is characterized by a severe, distressing unilateral head or face pain lasting from about 15 minutes to 3 hours. The localization is usually orbital or temporal, and the onset and offset are rapid. The pain is described as burning, piercing, throbbing or pulsing. The full syndrome includes conjunctival injection, forehead sweating, meiosis, ptosis, lacrimation, eyelid oedema and nasal congestion. The pain occurs once or more than once daily for weeks to months, with pain-free intervals of 2 weeks or more. Alcohol may precipitate an attack. During an attack, the patient will often pace around the room, grasping the affected eye in an attempt to relieve the pain. This is in contrast to the patient with migraine who will usually wish to lie still in a quiet, dark environment. Sumatriptan subcutaneously is the drug of choice in the treatment of acute attacks; inhalation of 100 per cent oxygen can also be very helpful. For short-term prophylaxis, oral corticosteroids, methysergide or ergotamine may be helpful, while for longer-term...
prophylaxis verapamil is a first-line treatment. About 10–20 per cent of patients do not have remissions (or remissions of less than 2 weeks) and are classified as having chronic cluster headache.

**Short-lasting unilateral neuralgiform headache attacks with conjunctival injection and tearing (SUNCT)**

This is another trigeminal autonomic cephalgia syndrome, distinguished from cluster headache by brief (average 50 seconds), frequent attacks (up to 30 per hour) with prominent autonomic features in the majority of patients. The treatment of SUNCT is challenging. Some response has been seen with carbamazepine, and more recently lamotrigine, which is a promising potential first-line treatment. Further data from randomized trials are awaited.

**Paroxysmal hemicrania**

This is also a short-duration unilateral headache syndrome with pain in the maxillary, orbital, frontal or temporal regions and associated autonomic features. The characteristic and distinguishing feature is the excellent response to indometacin treatment. Unlike cluster headache, only half of the patients tend to pace around when the pain occurs.

**VASCULAR DISORDERS**

A mild headache occurs at the onset of about one-quarter of ischaemic stroke or transient ischaemic attacks. It has been reported to be more common with vertebrobasilar territory events than with carotid territory events, and it is rarer still in lacunar syndromes. If head pain is lateralized and severe, and followed by fixed neurological deficit, arterial dissection must be considered. Carotid dissection typically causes unilateral pain localized to the face, frontal region or eye, whereas vertebral dissection may cause unilateral or bilateral occipital pain. Headache of raised intracranial pressure-type (i.e. headache worse on waking, exacerbated by coughing, sneezing or straining) may be a feature of cerebral venous sinus thrombosis, in which focal signs including seizures are often found. The usual presentation of venous sinus thrombosis is with a subacute raised pressure headache rather than an acute headache.

Sudden, severe headache is the most important symptom of subarachnoid haemorrhage, and it may be the only complaint in one-third of patients. It occurs at some stage in the illness in between 85 and 100 per cent of cases. The cardinal feature of this type of headache is its exceptionally rapid onset. It may be described as being ‘like a hammer blow to the head’, or ‘like an explosion in the head’, and most patients will volunteer that it is the worst headache that they have ever experienced. It reaches a maximum in a split second, or within a few seconds (conventionally the upper limit is taken to be about 1 minute). This is termed ‘thunderclap headache’ and, although subarachnoid haemorrhage is the most frequent serious cause, there are other causes, as listed in Box H.7. Other symptoms suggestive of subarachnoid haemorrhage include occipital localization of the pain, neck stiffness, nausea, vomiting, and exertion or Valsalva immediately preceding the headache.

Up to about 40 per cent of patients with aneurysmal subarachnoid hemorrhage report a history of a sentinel or warning headache, days to weeks prior to aneurysm rupture; these may be caused by small blood leaks into the subarachnoid space or aneurysm wall changes. A history of thunderclap headache makes computed tomographic (CT) scanning mandatory to seek evidence of acute subarachnoid blood, which appears as high attenuation (bright). If the CT scan is negative, a lumbar puncture must be performed to look for red cells or their breakdown products. The most common focal neurological sign in subarachnoid haemorrhage is of a lillrd nerve palsy.

**Reversible cerebral vasoconstriction syndrome (RCVS)** is a syndrome of thunderclap headache, often recurrent, which has confusingly been described by a number of different terms including: benign angiopathy of the central nervous system; migrainous vasospasm or ‘crash migraine’; Call-Fleming syndrome; and postpartum angiopathy. It is almost certainly under-recognized and under-diagnosed and may underlie some classical primary headache syndromes, e.g. benign thunderclap headache or coital cephalgia. It is likely that some individuals diagnosed with CNS vasculitis also have RCVS. RCVS is important to diagnose as it is often self-limiting without specific treatment.

Drug triggers for RCVS are common, including ergotamine, triptans, selective serotonin reuptake

**Box H.7 Causes of thunderclap headache**

- Subarachnoid haemorrhage, usually aneurysmal
- Reversible cerebral vasoconstriction syndromes
- Sentinel headache
- Cerebral venous thrombosis
- Cervical artery dissection
- Spontaneous intracranial hypotension
- Pituitary apoplexy
- Retroclival hematoma
- Ischemic stroke
- Acute hypertensive crisis
- Colloid cyst of the third ventricle
- Infections
- Primary thunderclap headache
Intraventricular haemorrhage bilateral visual loss. headache, classically in association with sudden can cause a similar sudden severe Pituitary apoplexy have similar efficacy for stroke prevention and good outcome. hematoma. Management is with antithrombotic flap, double lumen and intramural crescent-shaped dissecting aneurysm (pseudoaneurysm), intimal arterial stenosis or occlusion (‘rat’s tail sign’), Diagnostic imaging findings include a long, tapered and axial MRI of the neck with fat saturation protocol. for dissection include CT angiography, magnetic resonance angiography, conventional angiography, and cranial neuropathies. Useful diagnostic tests dissection include Horner syndrome, pulsatile tinnitus is described but is rare. Other (local) symptoms of stenosed dissected artery. Subarachnoid haemorrhage ischaemia, due to reduced perfusion through a abnormal dissected vessel lumen or haemodynamic ischaemia, due to reduced perfusion through a stenosed dissected artery. Subarachnoid haemorrhage is described but is rare. Other (local) symptoms of dissection include Horner syndrome, pulsatile tinnitus and cranial neuropathies. Useful diagnostic tests for dissection include CT angiography, magnetic resonance angiography, conventional angiography, and axial MRI of the neck with fat saturation protocol. Diagnostic imaging findings include a long, tapered arterial stenosis or occlusion (‘rat’s tail sign’), dissecting aneurysm (pseudoaneurysm), intimal flap, double lumen and intramural crescent-shaped hematoma. Management is with antithrombotic agents: antiplatelets and anticoagulants appear to have similar efficacy for stroke prevention and good outcome.

Pituitary apoplexy can cause a similar sudden severe headache, classically in association with sudden bilateral visual loss. Intraventricular haemorrhage can also mimic the headache of subarachnoid haemorrhage.

Primary intracerebral haemorrhage, especially in a peripheral lobar distribution, is preceded by headache in one-half of patients. Focal neurological deficit is almost always found.

It is rare for unruptured intracranial aneurysms to cause significant headache, but occasionally a large posterior communicating artery aneurysm may cause pain localized to behind the eye. Temporal arteritis must be remembered because it is a preventable cause of blindness. It usually occurs in patients aged over 50 years, and the prevalence increases with increasing age. Head or face pain is found at presentation in 50 per cent of cases. One-quarter of patients have systemic symptoms at presentation (arthralgia, myalgia or low-grade pyrexia). Pain in the temporal or masseter muscles on chewing (jaw claudication) is highly characteristic and virtually diagnostic. Blindness occurs due to involvement of the posterior ciliary artery, which supplies the optic disc. On examination, the temporal artery may be palpable and tender. The erythrocyte sedimentation rate is usually markedly elevated (up to 140 mm/h). The acute treatment is high-dose oral or intravenous corticosteroids.

Cerebral venous sinus thrombosis often presents with headache, but it is only of sudden onset in about 15 per cent of patients. Most patients will have symptoms and signs suggesting raised intracranial pressure, and about one-third will experience seizures or focal neurological signs (e.g. hemiparesis).

NON-VASCULAR INTRACRANIAL PATHOLOGY

Raised intracranial pressure

High intracranial CSF pressure causes a characteristic headache that has already been briefly mentioned. The headache is typically aching or throbbing, exacerbated by coughing, bending over or straining, worse in the morning, and it may cause wakening from sleep. It may be worse after exertion or on lying flat. As the headache worsens, vomiting, diplopia and papilloedema develop. The raised intracranial pressure may be due to obstruction of normal CSF flow by an intracranial tumour or by another mechanism affecting CSF dynamics. The first clinical priority if high intracranial pressure is suspected is to exclude an intracranial mass lesion by imaging – with CT or ideally magnetic resonance imaging (MRI) (which provides better quality images of the posterior fossa). If there is no mass lesion, raised pressure must be due to another mechanism (Box H.8).
Raised pressure may also be due to impaired CSF reabsorption; this may result from current or previous meningitis (acute, subacute or chronic, e.g. granulomatous or carcinomatous) or from subarachnoid haemorrhage; both of these processes block the arachnoid granulations. Thus, once a mass lesion is excluded by imaging, the CSF should be examined by lumbar puncture to seek red or white cells and to determine the protein concentration. If the CSF is normal and there is no mass lesion, and the patient has focal hemisphere signs or seizures, cerebral venous sinus thrombosis must be considered and excluded by either formal or magnetic resonance venography studies.

If CSF pressure is elevated but focal neurological signs are minimal or absent, and both imaging (including venography) and CSF analysis are normal, the patient has a syndrome of idiopathic intracranial hypertension (formerly called benign intracranial hypertension or pseudotumour cerebri). In its classical form, this syndrome occurs in women who are overweight and often with menstrual irregularities. As well as a raised pressure headache, patients will often complain of transient visual loss on changing posture (e.g. bending over), termed ‘visual obscurations’. Diplopia or unilateral facial numbness may also be present. A number of associations exist with this syndrome, including the use of tetracycline antibiotic, lead poisoning, vitamin A toxicity and metabolic disturbance (hypothyroidism).

Some conditions cause raised intracranial pressure by dramatically raising the CSF protein concentration (Guillain–Barré syndrome, spinal oligodendroglioma or systemic lupus erythematosus).

**Box H.8 Causes of raised intracranial pressure**

- Cerebral, dural or extradural mass causing impaired CSF flow
  - Tumour
  - Abscess
  - Haematoma
- Generalized brain swelling
  - Hypoxia
  - metabolic disturbance
  - Hypertensive encephalopathy
- Increase in cerebral venous pressure
  - Heart failure
  - Obstruction of superior mediastinal or jugular veins
  - Cerebral venous sinus thrombosis
- Obstruction to CSF resorption or flow
  - Meningitis (granulomatous, carcinomatous, haemorrhagic)
  - Subarachnoid haemorrhage
- Expansion of CSF volume
  - CSF secreting tumour (choroid plexus tumour) – rare
- Unknown mechanism
- Idiopathic intracranial hypertension

**Low CSF pressure headache**

Low CSF pressure can also cause a distinctive headache syndrome. This type of headache may follow lumbar puncture. The pain is not usually present on waking, is worse on sitting up or standing, and is rapidly relieved by lying flat. The presumed mechanism is of persistent CSF leakage through a dural tear. The management of post-lumbar puncture headache is bed rest, fluids and analgesia; it may take several weeks to resolve, and in severe cases epidural blood patches have been used. Sometimes, low-pressure headache can occur spontaneously or following head, neck or spinal trauma. In other cases, there may be an index Valsalva event such as coughing, straining, lifting, etc. In low-pressure headache, MRI with contrast demonstrates striking meningeal enhancement. The management is similar to that of post-lumbar puncture headache. Intravenous caffeine or theophylline has been effective in some cases.

**Intracranial infection**

**Meningitis** is inflammation of the pia and arachnoid, caused by bacterial, viral, fungal or other infections. The clinical syndrome of meningeal irritation must be recognized quickly so that appropriate treatment can be commenced. Untreated bacterial meningitis can be very rapidly fatal. The headache in acute meningeal irritation is of rapid onset (less than 48 hours) and severe. There are accompanying symptoms of photophobia, and drowsiness, vomiting, irritability and seizures may develop. The important physical signs are fever, neck stiffness, **Kernig’s sign** (pain and resistance when the examiner extends the knee with the hip fully flexed) and **Brudzinski’s sign** (the hips flex when the head is flexed forward towards the chest). Lumbar puncture must be performed rapidly to confirm the diagnosis and the organism, unless there is drowsiness, suspicion of raised intracranial pressure or focal neurological signs, in which case CT must be performed first, and empirical antibiotics commenced. The most common organisms in adults are **Meningococcus** and **Pneumococcus**, but in those who are immuno-compromised or elderly **Listeria**, fungi (including **Cryptococcus**) and tuberculosis must also be considered. In the immunocompromised, some of the typical features of meningitis, such as neck stiffness and a positive Kernig’s sign, may be absent. **Aseptic meningitis** may give an identical clinical syndrome, but no organisms are identified in the CSF. **Chronic meningitis** may be due to granulomatous or carcinomatous meningeal irritation, and it will have...
HEADACHE

a less acute onset and often associated cranial nerve palsies. In patients with HIV infection and advanced disease, headache may be due to infection with Toxoplasma gondii. This characteristically produces multiple ring-enhancing intracerebral lesions on CT or MRI scans; the diagnosis is confirmed by a response to empirical therapy, or much less often by brain biopsy.

Encephalitis is inflammation of the brain parenchyma, although a variable degree of meningeal involvement usually occurs as well. The patient will often have similar symptoms to those of meningitis, but there may be a subacute history of personality change, and seizures with focal signs (e.g. hemiparesis) are common. Viruses (e.g. herpes simplex, Coxsackie, echovirus and rabies) are the most common cause and, if the diagnosis is suspected, treatment with aciclovir should be started empirically.

Cerebral tumour

Although headache is an early symptom in about one-third of brain tumours, the vast majority of headaches are not due to tumours. There are no specific features, although the features of raised intracranial pressure (see above) may develop if the tumour is impeding the CSF circulation. Tumours may cause headache without raised CSF pressure; the presumed mechanism is the distension of local, pain-sensitive structures (blood vessels and dura). If lateralized headache is present, it is very often on the side of the tumour. It is self-evident that a tumour may produce focal neurological symptoms and signs depending on its location, although some present with only subtle cognitive disturbance. New-onset seizures in an adult should always raise the suspicion of cerebral tumour. Any headache of recent onset with focal neurological symptoms and signs (especially seizures), or papilloedema, requires urgent neuroimaging. Pupillary abnormalities and reduced consciousness level are usually late symptoms reflecting tentorial herniation or rapidly rising intracranial pressure.

Headache associated with systemic infection

Many infectious illnesses are associated with headache, for example viral infections. There will be other symptoms to indicate the systemic nature of the illness.

Specific headache syndromes

It is helpful to be able to recognize these syndromes, as doing so will allow a firm diagnosis to be made, and reassurance or appropriate treatment to be offered.

• Idiopathic stabbing headache: This describes sharp, stabbing pain in the head that may occur once or in recurrent volleys. The pain is usually in the first trigeminal division, lasts only a split second and recurs irregularly (hours to days apart). It is more common in women and tends to be spontaneous rather than triggered. There are no autonomic features, and indomethacin is an effective treatment.

• Cold stimulus headache: This condition, also called ‘ice-cream headache’, is reported to occur in about one-third of the population. Cold stimulation of the pharynx or palate induces pain shortly afterwards (10–20 seconds) that is often lateralized and is of short duration (seconds), but it can be longer. The pain may be referred to the forehead or temple by the trigeminal nerve, or to the ears by the glossopharyngeal nerve. In recent studies, ice-cream headache has been reported to be less common in patients with migraine than in the general population.

• Benign cough and exertional headache: Some patients report severe, transient head pain on coughing, sneezing, laughing vigorously or lifting heavy weights, bending over or straining. The pain is typically frontal or occipital, follows the action by 1–2 seconds, and lasts for only a few seconds. It may be described as ‘explosive’ or ‘bursting’. It often occurs recurrently for months to a year or two, and then remits. This is a benign syndrome, but serious pathology can cause identical symptoms, for example posterior fossa tumours, so investigations may be needed. Runners and other athletes often get headaches on exertion. These headaches may have associated features suggesting that they are a form of migraine. Exertional and cough headaches may respond to indomethacin.

Headache associated with head trauma

Headaches affect most symptomatic patients after mild head injury. Post-traumatic headaches usually begin hours or days after the injury, but they may be delayed for weeks. Headaches may paradoxically occur more often and for longer in patients with mild compared with severe trauma. Commonly associated symptoms include dizziness, irritability, lack of concentration and intolerance to alcohol. Vertigo, hearing disturbance,
apathy and tiredness may also occur as part of the ‘post-head injury syndrome’. Headaches after trauma may be localized or diffuse, episodic or daily. They most often resemble chronic tension-type headaches but may be migrainous. Neuralgic or trigeminal autonomic cephalgia-type pains less commonly occur. If the dura has been damaged, low-pressure headaches may result.

Headache associated with substance use
Some substances reliably induce headaches during acute use in some individuals. Nitrates, found in preserved, processed or cured meats, cause so-called ‘hot-dog headache’ which is usually bitemporal and associated with facial flushing. Monosodium glutamate causes what has been termed ‘Chinese restaurant headache’, where associated features include chest tightness, and burning in the head and upper trunk. Carbon monoxide is an important cause of headache, but there will usually be associated features including reduced consciousness level. Hyperbaric oxygen treatment can be rapidly effective. Alcohol-induced headache occurs within hours of ingestion (unlike typical ‘hangover’ headache, which is delayed) and is most commonly migrainous. Chocolate and cheese have long been thought to be important triggers for migraine, but they can also cause headaches in non-migraineurs.

Headache associated with substance misuse
Headache from substance misuse can occur with central nervous system stimulants (amphetamine, cocaine and designer drugs), barbiturates and sedatives, or opiates.

The headaches may be associated with acute or ‘binge’ use or may occur during withdrawal from these substances.

HEART, DISPLACED IMPULSE
Gerry Carr-White

The position of the cardiac apex is defined as the furthest outward and downward point at which a cardiac impulse can be felt in the supine, unrotated patient. This point usually lies in the fifth or sixth left intercostal space in the mid-clavicular line. Displacement of the apex may be due to a congenitally abnormal position of the heart, to cardiac enlargement, or to cardiac displacement resulting from chest wall or intrathoracic abnormalities.

Congenital dextrocardia may involve simple mirror-image inversion of the heart and other viscera (situs inversus) or may be associated with a variety of complex congenital cardiac defects. In adults, simple dextrocardia without associated defects is more common and is benign (Fig. H.8).

Displacement of the cardiac apex resulting from cardiac enlargement can usually be suspected on the basis of the patient’s symptoms of heart disease, or the presence of murmurs or abnormal heart sounds. Confirmation can be made by chest radiography, electrocardiography and echocardiography. The differential diagnosis is discussed more elsewhere (see HEART, ENLARGEMENT OF, p. 266).

Displacement of the cardiac apex from abnormalities of the chest wall is usually apparent on inspection of the patient; these abnormalities include pectus excavatum (funnel chest), pectus carinatum (pigeon chest), kyphoscoliosis and thoracoplasty.

An exception is sometimes a ‘compensated’ thoracic kyphoscoliosis, which may be more easily appreciated on a chest X-ray. Conversely, pectus excavatum is obvious clinically but is sometimes missed on cursory inspection of a posteroanterior chest radiograph.

Displacements of the cardiac apex as a result of intrathoracic abnormalities are listed in Box H.9, according to the organ or tissue involved. Pulmonary or diaphragmatic causes are common, the remainder are rare.

LYMPHOID TISSUE AND THYMUS
Enlargement of thoracic lymph nodes is common but seldom causes cardiac displacement. Rarely, enlargement of para-aortic nodes or aberrant thymus tissues may do so.
HEART, ENLARGEMENT OF

Box H.9 Displacement of the cardiac apex

- Lung and pleura
  - Pleural effusion
  - Pneumonectomy
- Pneumothorax
- Tension pneumothorax
- Diaphragm
  - Congenital diaphragmatic hernia
  - Hiatus hernia
- Pericardium
  - Clearwater cyst of pericardium
  - Pericardial tumour
- Oesophagus
  - Achalasia of the cardia/megaesophagus
  - Foregut duplication cyst
- Aorta
  - Aneurysm of descending aorta

The differential diagnosis can often be clarified by radiography in the posteroanterior and lateral projections, with the help of a barium swallow to outline the oesophagus. In more difficult cases, computed tomography scanning and echocardiography – particularly transoesophageal echocardiography – are useful.

HEART, ENLARGEMENT OF

Melvin Lobo

Cardiac enlargement may be detected or suspected on the basis of clinical examination (see HEART, DISPLACED IMPULSE, p. 265), a chest radiograph or an electrocardiogram. However, enlargement of the cardiac chambers can only be definitively shown by echocardiography or other techniques such as cardiac magnetic resonance scanning. Hypertrophy (usually of the left ventricle) refers to increased thickness of the walls and may or may not be associated with enlargement of the chambers. Myocardial mass refers to the total weight of the heart (usually assessed non-invasively with echo or MRI). When considering a patient with apparent cardiac enlargement, it is necessary to decide: (i) whether the enlargement is genuine or spurious; (ii) if genuine, whether it is physiological or pathological; and (iii) if pathological, whether it is due primarily to a myocardial disorder or a haemodynamic lesion.

THE IDENTIFICATION OF GENUINE CARDIAC ENLARGEMENT

The clinical diagnosis of cardiac enlargement is usually made because of displacement of the cardiac impulse. With left ventricular hypertrophy due to hypertension, aortic stenosis or hypertrophic cardiomyopathy, the apex may not be displaced but has a heaving, sustained character quite different from the normal.

Radiographic cardiac enlargement is diagnosed when the cardiothoracic ratio is greater than 0.5 in a posteroanterior chest radiograph, with a tube-to-film distance of at least 2 m, taken in full inspiration. The cardiothoracic ratio is the ratio of the widest part of the cardiac shadow to the widest part of the lung fields. A spurious impression of cardiomegaly is caused by anteroposterior projections, portable apparatus with a reduced tube–film distance, and a poor inspiratory effort. In pectus excavation (funnel chest), a false impression of cardiac enlargement may be obtained from a posteroanterior film, but a lateral film will clarify the issue.

Radiological enlargement of the cardiac shadow may be due to a pericardial effusion. This is more likely if the enlargement is rapid (and previous films are available for comparison). Clinically, there will usually be elevation of the jugular venous pressure; there may be a pericardial friction rub, and an increased area of cardiac dullness. The electrocardiogram (ECG) usually shows low voltages, and there may be electrical alternans (alternating large and small QRS complexes in the same lead). Echocardiography is diagnostic.

An electrocardiographic diagnosis of cardiac enlargement is usually inferred by general abnormalities such as an abnormal axis, pathological Q waves or T wave inversion or voltage criteria for left ventricular hypertrophy (sum of the S wave in lead V2 and the R wave in V5 of greater than 3.5 mV is one of the criteria for left ventricular hypertrophy, but this criterion is sometimes met in fit muscular young men with thin chest walls).

THE IDENTIFICATION OF A PATHOLOGICAL PROCESS

Athletes tend to have enlarged hearts (often both left and right ventricles are enlarged and there may be mild atrial enlargement), a slow resting pulse rate, and often a soft ejection systolic murmur related to a large resting stroke volume. In ‘duration’ sports (e.g. running and swimming), there is chamber enlargement without disproportionate hypertrophy of the ventricular walls. In ‘power’ sports (e.g. weightlifting), there may also be ventricular hypertrophy. If left ventricular hypertrophy is marked on electrocardiography or echocardiography, it is worth asking about – and warning against – the abuse of anabolic steroids.

Cardiac enlargement is sometimes found in patients with congenital heart block, as an adaptive response to a slow pulse rate. It is usually present in acromegaly, sometimes as part of a general process of soft-tissue hypertrophy but sometimes as a feature of acromegalic heart muscle disease.
THE DISTINCTION BETWEEN MYOCARDIAL AND HAEMODYNAMIC DISORDERS

Cardiac enlargement, without a murmur but with an added third or fourth heart sound, is more likely to be due to a myocardial disorder and, with a murmur, to a haemodynamic lesion; the distinction is not absolute, however. The best diagnostic tool is duplex ultrasound (echocardiography plus Doppler cardiography), which gives information about both valve performance and ventricular function.

Myocardial problems

The principal heart muscle problems (in order of frequency, in ‘Western’ practice) are ischaemic heart disease, dilated cardiomyopathy, hypertrophic cardiomyopathy and the ‘specific heart muscle diseases’, which mimic dilated cardiomyopathy. Cardiac enlargement due to right ventricular enlargement is often seen in patients with congenital heart disease (such as tetralogy of Fallot and large ASDs). Cardiac enlargement due to atrial enlargement may be seen in restrictive cardiomyopathies, mitral stenosis and sometimes chronic atrial fibrillation.

Ischaemic heart disease

Myocardial infarction is the commonest cause of dilatation of the left ventricle. This is more frequent after anterior infarction. The cardiac impulse is displaced and may be ‘hypokinetic’. The ECG may show evidence of old infarction, sometimes with persistent ST segment elevation suggesting an aneurysm. The chest X-ray usually shows cardiomegaly (Fig. H.9), occasionally with calcification in the aneurysm. Echocardiography is diagnostic. Ischaemic heart disease can also cause global left ventricular dilatation that is indistinguishable clinically from a dilated cardiomyopathy, although symptoms of angina or a history of myocardial infarction may provide clues.

Dilated cardiomyopathy

Dilated cardiomyopathy is left ventricular dilatation and dysfunction in the absence of coronary disease, valve disease or hypertension, and a cause can be found in about 50 per cent of cases. In those without a clear cause, there is often evidence that they are genetic, typically inherited in an autosomal dominant manner. Myocarditis refers to an acute inflammatory myocardial process (due to a variety of causes such as viral infections and autoimmune conditions) that often leads on to a dilated cardiomyopathy in the chronic phase. A list of disorders leading to a dilated cardiomyopathy is shown below (Box H.10). (Note that this is not an exhaustive list of specific heart muscle diseases, but emphasizes those which may present with cardiac enlargement.)

The heart is clinically and radiologically enlarged, and there may be features of cardiac failure, such as a tachycardia and often a gallop rhythm with a loud third heart sound. The apex is displaced, and there may be soft systolic murmurs from functional mitral or tricuspid regurgitation. The ECG is usually abnormal, but the changes are non-specific. Echocardiography shows a characteristic pattern with dilatation of the left ventricle and often the other chambers as well and reduced left ventricular contraction. Myocardial biopsy has a limited role, largely in excluding myocarditis or a specific cause. The prognosis is best when there is a remediable cause such as alcohol abuse, or where acute myocarditis is followed by spontaneous recovery.

Acute viral myocarditis is a common cause of a dilated cardiomyopathy pattern in younger adults. There is often a preceding acute febrile illness or respiratory tract infection. Many viruses have been implicated, but the most consistent link is with the Coxsackie B group. The vast majority of cases of myocarditis probably go unrecognized, but the prognosis in cases that present with cardiac failure is poor.

Alcoholic heart disease is another common cause of the specific heart muscle diseases in middle-aged Western males, and is often very reversible on cessation of alcohol. Amyloid heart disease is unusual among the specific heart muscle diseases in that ventricular dilatation is not a prominent feature, although ventricular function is impaired.

Figure H.9 Chest radiograph showing an enlarged heart (cardiomegaly).
Hypertrophic cardiomyopathy

Hypertrophic cardiomyopathy is characterized by increased left ventricular wall thickness (>15 mm) in the absence of abnormal loading conditions such as aortic stenosis or hypertension. It occurs in 1 in 500 of the population and is most commonly due to genetic mutations in genes encoding the sarcomere proteins. The disease is usually inherited as an autosomal dominant trait and tends to present in adolescence or early twenties, though many patients are asymptomatic and unaware of their condition. Symptoms include breathlessness, chest pain and blackouts. There is an increased risk of sudden death, occurring in about 1 per cent of patients a year. Factors that increase the risk are syncope, a family history of early sudden death, non-sustained ventricular tachycardia on monitoring, left ventricular wall thickness above 30 mm and an abnormal blood pressure response on exertion.

Typically, the left ventricle is hypertrophied but not dilated, though in about 10 per cent of cases the left ventricle does dilate. The apex is displaced and often has a ‘double beat’ character from a palpable atrial impulse, as well as the sustained lift of ventricular hypertrophy. In about a third of patients, there is left ventricular outflow tract obstruction due to systolic anterior motion of the mitral valve towards the thickened septum. In these cases, there is an ejection systolic murmur at the left sternal edge, which becomes louder sitting up and during a Valsalva manoeuvre. There may be reversed splitting of the second heart sound, and the peripheral pulses are ‘jerky’. The ECG is abnormal, typically showing left ventricular hypertrophy with marked T wave changes, and sometimes pathological Q waves in the absence of prior infarction. Echocardiography is diagnostic. In some patients with hypertrophic cardiomyopathy, there are underlying disorders (so-called ‘phenocopies’). Examples include patients with the hereditary neurological disorder of Friedreich's ataxia, the metabolic disorder Fabry's disease and patients with Noonan's syndrome.

Haemodynamic problems

Haemodynamic problems causing cardiac enlargement can be divided into those causing a volume load and those causing a pressure load (Boxes H.11 and H.12). The initial response to a volume load is an increase in stroke volume, with enlargement of the cardiac chambers concerned, but with preservation of vigorous contraction. The response to a pressure load is hypertrophy, without dilatation, of the chambers involved. If either a volume or a pressure load persists beyond the capacity of the heart to compensate, the chambers will dilate, and contraction will become impaired. Chronic failure of the ‘left side’ of the heart, from mitral valve disease, or left ventricular impairment, may be accompanied by reactive pulmonary vasoconstriction and secondary pulmonary
hypertension, with ensuing dilatation of the right heart chambers as well.

Systemic hypertension is by far the most common ‘pressure load’ cause of cardiac enlargement. Most cases are due to ‘essential’ hypertension, but renal and endocrine causes should be remembered. Aortic coarctation causes delayed femoral pulses, and there is often a systolic murmur. A phaeochromocytoma can cause severe, but intermittent, hypertension.

Murmurs should lead to the diagnosis of aortic or pulmonary stenosis. Aortic stenosis is frequently severe even when there is little or no radiological cardiac enlargement, but the presence of poststenotic dilatation of the ascending aorta may give a clue.

The diagnosis of pulmonary hypertension is often missed. The circumstances may be suggestive, and clinically the combination of a pronounced right parasternal impulse with changes of right ventricular hypertrophy in the ECG should lead it to be suspected.

Mitral and aortic regurgitation are detected through their distinctive murmurs. Selective enlargement of the different cardiac chambers may give a characteristic radiographic appearance. However, in some cases, the radiographic appearance is non-specific and further aid must be sought from ultrasound scanning.

### HEART, MURMURS IN

Gerry Carr-White

A heart murmur is the audible sign of turbulent blood flow within the heart. The site of maximum intensity of the murmur should be noted, together with the direction in which it radiates – as a rule, this is the direction in which the turbulent blood is flowing.

Other clues such as cardiac enlargement or abnormal pulsation in the precordium or neck vessels should also be sought. A thrill is a palpable vibration, and, for practical purposes, precordial thrills can be regarded as ‘palpable’ murmurs. High-pitched vibration is usually more easily heard than felt, and the significance of a thrill is simply that of a loud murmur. With low-frequency vibrations, some observers claim to find a thrill easier to feel than a low-pitched murmur is to hear.

Heart murmurs can be classified as either systolic or diastolic (the continuous murmur of persistent ductus arteriosus is, strictly speaking, exocardiac), although a single lesion can sometimes cause both. Systolic murmurs accompany the upstroke of the carotid pulse; diastolic murmurs precede or follow it.

### SYSTOLIC MURMURS

These are further classified as ejection systolic murmurs, which increase to a crescendo in mid-systole and then die away before the second sound, and pansystolic murmurs, which remain at a more constant amplitude throughout systole extending to the heart sounds. Late systolic murmurs are a variant of the latter, appearing in mid or late systole and continuing to the second sound (Fig. H.10).

### EJECTION SYSTOLIC MURMURS

The causes of these are listed in Box H.13.
Innocent systolic murmurs
An ejection systolic murmur is commonly heard in fit children or young adults, particularly after exercise or vasodilatation, for example during pregnancy. Characteristically, the murmur is soft, best heard in the second left intercostal space, and becomes louder on inspiration; the second sound is normally split (see HEART SOUNDS, p. 273), and there are no other abnormal signs. Further investigation of these patients is almost always negative, and the long-term prognosis is excellent.

Aortic and pulmonary stenosis
The murmurs of mild aortic or pulmonary stenosis differ from innocent murmurs, principally in that they tend to be louder and the murmur is often preceded by an ejection click. Most patients with mild congenital aortic stenosis have bicuspid valves. Eventually, the abnormal valve tends to calcify and a mild stenosis may become severe. In elderly patients, a soft ejection systolic murmur may be heard beneath the left scapula; other signs such as delayed femoral pulses are also present. In young patients, the murmurs of severe pulmonary or aortic stenosis tend to be loud and harsh, but loudness is a poor guide to the severity of aortic stenosis in the elderly or when the cardiac output is low. Other features of severe aortic stenosis include clinical evidence of left ventricular hypertrophy and a slow-rising carotid pulse. In severe pulmonary stenosis, the second sound is widely split. Two-dimensional echocardiography plus Doppler studies is the most useful investigation.

Flow murmurs
Ejection systolic murmurs may be due to turbulence caused by a normal blood flow through a stenotic valve orifice, as in aortic stenosis, or by an increased blood flow through a non-stenotic valve. The latter mechanism accounts for most innocent murmurs, and for the pulmonary ejection systolic murmur of atrial septal defect, in which the flow through the pulmonary valve orifice may be increased two- or threefold above normal. These patients have an ‘active’ feel to the precordium, there is fixed splitting of the second sound, and there may be a tricuspid diastolic flow murmur. Similarly, patients with marked aortic regurgitation may have a systolic murmur, despite the absence of anatomical stenosis, due to the increases stroke volume.

Hypertrophic cardiomyopathy
Between 30 and 40 per cent of these patients will have left ventricular outflow tract (LVOT) obstruction. They usually have an ejection systolic murmur, clinical evidence of left ventricular hypertrophy, and often a characteristically ‘jerky’ pulse. In these patients, the murmur paradoxically tends to become softer in inspiration and louder during expiration or in a Valsalva manoeuvre. Echocardiography is usually diagnostic.

PANSYSTOLIC MURMURS
The causes of such murmurs are listed in Box H.14.

Mitral regurgitation
The murmur of mitral regurgitation is usually loudest at the apex and radiates to the axilla. Loudness is some indication of severity, but there are exceptions. Torrential mitral regurgitation from papillary muscle rupture may be almost silent, while some patients with mitral valve prolapse and haemodynamically mild regurgitation may occasionally emit exceedingly loud ‘honks’ or ‘whoops’ that are audible across the room. The severity can be assessed from the patient’s general condition, cardiac enlargement, the presence of cardiac
failure and the radiographic appearances, as well as from the murmur. The term ‘mitral valve prolapse’ is used, confusingly, to describe either a diverse group of causes of non-rheumatic mitral regurgitation that have in common excessive elongation of the chordae tendineae, or a clinical syndrome that includes both the consequences of mitral regurgitation and also other features such as a predisposition to arrhythmias and, possibly, sudden death. Echocardiography is sometimes able to distinguish patients who simply have thin, stretched chordae tendineae (pellucid valve syndrome) from those with thickened, redundant, myxomatous valve tissue. It is in the latter group that most of the ‘extra-valvar’ events occur, though this does remain controversial.

Clinically, patients with mitral valve prolapse are sometimes indistinguishable from those with other causes of regurgitation. The murmur of prolapse tends, however, to be mid- or late-systolic rather than pansystolic, at least in the early stages, and there may be one or more mid-systolic ‘clicks’ due to tensing of the chordae.

**Congenital ventricular septal defect**

This is the most common cause of a pansystolic murmur in children and young adults. Large ventricular septal defects (VSDs) can cause cardiac failure within the first 3 months of life and, if untreated, sometimes induce reactive pulmonary hypertension with diminution (and eventual reversal) of shunt blood flow, cyanosis and disappearance of the murmur. Most pansystolic murmurs in older children are due to small defects with a loud murmur but a small shunt (maladie de Roger). The murmur is heard all over the precordium but maximally at the upper left sternal edge. Echocardiography is the investigation of choice. An accurate knowledge of the size and position of a defect helps to predict whether it is likely to close spontaneously.

**Acquired ventricular septal defect**

In adults, this may follow stab wounds or, more commonly, acute myocardial infarction. Clinical distinction between post-infarction septal defect and mitral regurgitation is difficult, but ultrasound is diagnostic.

**Tricuspid regurgitation**

This may be primary, from rheumatic heart disease or endocarditis, or secondary to right ventricular dilatation in response to pulmonary hypertension. The murmur is loudest at the lower left sternal edge, and is accompanied by pathognomonic prominent ‘v’ waves in the jugular venous pulse. This lesion is seen much more often in conjunction with other cardiac lesions than on its own.

**Pericardial ‘crunch’**

Patients with acute rupture of the oesophagus may develop a noise in the chest, synchronous with the heartbeat, that is virtually indistinguishable from a loud pansystolic murmur. It is presumably a consequence of surgical emphysema in the mediastinum being ‘crunched’ with each systole.

**DIASTOLIC MURMURS**

These can be divided into early diastolic and mid diastolic. Early diastolic murmurs are usually soft, decrescendo, high-pitched – ‘like the letter R whispered’ – and best heard with the diaphragm of the stethoscope. They immediately follow the second heart sound, or the downstroke of the carotid pulse, and they tend to be best heard at the left (sometimes right) sternal border with the patient sitting up, leaning forward and breathing out. Mid-diastolic murmurs are usually low-pitched, rumbling and best heard with the stethoscope bell (Fig. H.11).

There are only two important causes of early diastolic murmurs – aortic and pulmonary regurgitation. The causes of these lesions are listed in Box H.15.

**Box H.14 Causes of pansystolic murmurs**

<table>
<thead>
<tr>
<th>Most common</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Mitral regurgitation (in adults)</td>
</tr>
<tr>
<td>– Rheumatic</td>
</tr>
<tr>
<td>– Mitral valve prolapse</td>
</tr>
<tr>
<td>– Ischaemic</td>
</tr>
<tr>
<td>– Due to endocarditis</td>
</tr>
<tr>
<td>– Secondary to dilatation of mitral annulus</td>
</tr>
<tr>
<td>• Ventricular septal defect (in children)</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Common</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Tricuspid regurgitation</td>
</tr>
<tr>
<td>– Secondary</td>
</tr>
<tr>
<td>– Rheumatic</td>
</tr>
<tr>
<td>– Due to endocarditis</td>
</tr>
<tr>
<td>• Post-infarction ventricular septal defect</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Uncommon</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Pneumomediastium ‘pseudomurmur’ or ‘pericardial crunch’</td>
</tr>
</tbody>
</table>
Aortic regurgitation
Unless this lesion is mild, it is usually accompanied by a wide pulse pressure and a collapsing arterial pulse. Features such as exaggerated carotid pulsation (Corrigan’s sign), ‘pistol shot’ femoral bruits and capillary pulsation (Quincke’s sign) are ‘makeweights’ to be sought when the diagnosis is already firm. The murmur of acute aortic regurgitation, as in acute endocarditis, is similar in timing to the more usual early diastolic murmur but may be quite different in quality, being loud and harsh and apt to be confused with a systolic murmur, unless it is timed against the pulse.

Pulmonary regurgitation
This is usually secondary to pulmonary hypertension or in patients with complex congenital heart disease; the murmur of pulmonary regurgitation secondary to chronic rheumatoid mitral stenosis was described by Graham Steell. Pulmonary regurgitation may also be due to a damaged pulmonary valve resulting from a previous valvotomy or balloon valvuloplasty; in the absence of pulmonary hypertension, pulmonary regurgitation seems to be very well tolerated.

Distinction from aortic regurgitation is usually easy from the clinical circumstances and the absence of other features of aortic regurgitation. Confirmation is with echocardiography or cardiac MRI.

MID-DIASTOLIC MURMURS
The causes of these murmurs are listed in Box H.16.

Mitrail stenosis
This is by far the most common cause of a mid-diastolic murmur, which is best heard at the cardiac apex. There is often an audible or palpable opening snap in addition. The murmur of mitral stenosis is often accentuated just before the first sound (presystolic accentuation). This is usually only recognized in sinus rhythm, and attributed to an increased flow during atrial systole.

Carey Coombs murmur
A soft, low-pitched mid-diastolic murmur is sometimes heard in acute rheumatic fever, and it is named after the physician who first described it. It is not due to mitral stenosis but probably represents turbulence from minute vegetations on the mitral valve surface and from stiffening of the valve itself.

Austin Flint murmur
This murmur is due to fluttering of the anterior leaflet of the mitral valve, as a result of turbulence caused by aortic regurgitation. There is invariably an early diastolic murmur of aortic regurgitation, and the opening snap of mitral stenosis is absent. Echocardiography is diagnostic.

Mid-diastolic flow murmurs
The mid-diastolic murmur of increased tricuspid flow in atrial septal defect is usually heard only in children and adolescents. It tends to be higher pitched than the murmur of mitral stenosis and is best heard at the lower left sternal edge. A similar murmur may be heard at the apex in patients with a large VSD or persistent ductus arteriosus.

A box lists causes of early diastolic murmurs:

<table>
<thead>
<tr>
<th>Causes of early diastolic murmurs</th>
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</thead>
<tbody>
<tr>
<td>Aortic regurgitation due to</td>
</tr>
<tr>
<td>Most common</td>
</tr>
<tr>
<td>1. Bicuspid aortic valve</td>
</tr>
<tr>
<td>2. Chronic rheumatic heart disease</td>
</tr>
<tr>
<td>Common</td>
</tr>
<tr>
<td>1. Chronic hypertension</td>
</tr>
<tr>
<td>2. Dissecting aneurysm of aorta</td>
</tr>
<tr>
<td>3. Infective endocarditis</td>
</tr>
<tr>
<td>4. Failing aortic prosthesis</td>
</tr>
<tr>
<td>Uncommon</td>
</tr>
<tr>
<td>1. Acute rheumatic fever</td>
</tr>
<tr>
<td>2. Rheumatoid heart disease</td>
</tr>
<tr>
<td>3. Ankylosing spondylitis</td>
</tr>
<tr>
<td>4. Reiter’s syndrome</td>
</tr>
<tr>
<td>5. Chronic renal failure</td>
</tr>
<tr>
<td>Pulmonary regurgitation due to</td>
</tr>
<tr>
<td>Common</td>
</tr>
<tr>
<td>1. Congenital heart disease</td>
</tr>
<tr>
<td>2. Secondary pulmonary hypertension (e.g. in mitral valve disease)</td>
</tr>
<tr>
<td>3. Following pulmonary valvotomy or valvuloplasty</td>
</tr>
<tr>
<td>Uncommon</td>
</tr>
<tr>
<td>1. Primary pulmonary hypertension</td>
</tr>
<tr>
<td>2. Thromboembolic pulmonary hypertension</td>
</tr>
<tr>
<td>3. Eisenmenger’s syndrome</td>
</tr>
</tbody>
</table>

A box lists causes of mid-diastolic murmurs:

<table>
<thead>
<tr>
<th>Causes of mid-diastolic murmurs</th>
</tr>
</thead>
<tbody>
<tr>
<td>Common</td>
</tr>
<tr>
<td>1. Rheumatic mitral stenosis</td>
</tr>
<tr>
<td>Uncommon</td>
</tr>
<tr>
<td>1. Austin Flint murmur in aortic regurgitation</td>
</tr>
<tr>
<td>2. Tricuspid flow murmur in atrial septal defect</td>
</tr>
<tr>
<td>3. Tricuspid stenosis</td>
</tr>
<tr>
<td>4. Carey Coombs murmur in acute rheumatic fever</td>
</tr>
<tr>
<td>5. Left or right atrial myxoma</td>
</tr>
<tr>
<td>6. Mitral flow murmur in ventricular septal defect or persistent ductus arteriosus with large shunt</td>
</tr>
</tbody>
</table>
**Tricuspid stenosis**

The murmurs of tricuspid stenosis are similar to those of mitral stenosis, but the condition is much less common. The jugular venous pressure is elevated and the venous pulse has a characteristic waveform (see NECK, ENGORGED VEINS IN, p. 449).

**Atrial myxoma**

Left atrial myxomas are more common than right, although both are rare. They may cause murmurs mimicking mitral stenosis but, often, the murmurs vary with posture as the myxoma prolapses in or out of the valve orifice. The first sound is loud, and there may be an extra sound, called a ‘tumour plop’, from movement of the myxoma.

**HEART SOUNDS**

**Gerry Carr-White**

Tradition describes four heart sounds. The first and second are almost always audible; the third and fourth occur only in specific circumstances. There is also a medley of added sounds, mainly described as ‘clicks’, ‘snaps’ or ‘plops’, which can be heard in patients with particular conditions – these are distinguished from murmurs and rubs (see HEART, MURMURS IN, p. 269; RUB, PERICARDIAL, p. 593) by their short duration.

**FIRST SOUND**

The first heart sound is due to the closure of the mitral and tricuspid valves. Closure is normally simultaneous, but occasionally one valve closes slightly before the other, causing splitting of the first sound. This may be associated with right bundle-branch block, but it is seldom of clinical importance.

The first sound is usually readily recognized as the ‘lub’ in the traditional ‘lub-dup’ cadence of heart sounds. It is the sound that immediately precedes the upstroke of the carotid pulse. A loud first heart sound is most commonly associated with a hyperdynamic circulation (e.g. in pregnancy, febrile illness or thyrotoxicosis). Echocardiography has shown that the mitral cusps are still wide apart at the onset of systole as a consequence of the atrial augmentation of ventricular filling, and it is this, together with the increased rate of rise of ventricular pressure (dp/dt), as a result of sympathetic stimulation and a low peripheral resistance, which increase the force of valve closure. Mitral stenosis is the other common cause of an abnormally loud first sound. In rheumatic mitral stenosis, the mitral cusps are fused together laterally to form a diaphragm, which bulges into the left ventricle during diastole, and is propelled sharply back towards the left atrium in systole. The latter movement produces a loud, ringing first sound, often palpable as well as audible. A loud first sound is best heard in patients with a stenosed but still mobile valve in sinus rhythm. Progressive calcification restricts valve movement, and the sound gets quieter. A loud first sound alone does not make the diagnosis of mitral stenosis, but should prompt a careful search for a mid-diastolic murmur.

A much rarer cause of a loud first heart sound is a left atrial myxoma when it partially obstructs the mitral valve. Tricuspid stenosis may also cause a loud first sound, but this is rare. Apparently increased loudness of the first heart sound may simply be due to a relative lack of soft tissue between heart and stethoscope, as in fit, thin young subjects or patients who have had a mastectomy. Conversely, obesity or emphysema may muffle the heart sounds.

Varying intensity of the first sound occurs in three common conditions: atrial fibrillation, extra systoles, and complete heart block. In atrial fibrillation, the varying length of diastole causes the mitral cusps to be in varying positions at the onset of systole, and ventricular stroke volume also varies with the length of diastole. To some extent, these effects cancel out, so the variation in first sound intensity may be less than in complete heart block, where stroke volume is relatively constant but where the varying relationship of atrial to ventricular systole causes the position of the mitral cusps to vary from beat to beat at the onset of systole. With extra systoles, the fast sound of the premature beat is invariably softer. Coupled extra systoles cause a characteristic cadence of a loud first, normal second sound, soft first and second sound, pause. This is often misinterpreted because the premature beat gives no palpable pulse, and all the sounds are ascribed to a single cardiac cycle.

An abnormally quiet first heart sound, unless an artefact of obesity or emphysema, is usually due to a reduced cardiac output. In left ventricular failure, a small increase in left ventricular volume during diastole causes a large rise in pressure, and echocardiography shows that the mitral cusps have virtually drifted together before the onset of systole. In these circumstances, a quiet first heart sound frequently accompanies a third or fourth sound, as described below.

A sudden diminution in the intensity of the first sound may occur in acute mitral regurgitation, when it will be associated with the appearance of a pansystolic murmur. Even more rarely, an endocarditic vegetation or an atrial myxoma (see below) may interfere with mitral valve closure, and cause a sudden reduction in the first sound.
The changes in intensity of the first heart sound can be summarized as follows:

- Loud
- Thin chest wall
- Hyperdynamic circulation
- Mitral stenosis
- Atrial myxoma
- Soft
- Obesity/emphysema
- Cardiac failure
- Acute mitral regurgitation
- Endocarditis/myxoma
- Large pericardial effusion
- Variable
- Atrial fibrillation
- Extrasystoles
- Complete heart block
- Atrial myxoma

SECOND SOUND

The second heart sound is due to closure of the aortic and pulmonary valves. In expiration, their closure is normally synchronous, but in inspiration, there is a tendency for the aortic valve to close slightly earlier, the negative intrathoracic pressure causing blood to pool in the pulmonary veins, and the pulmonary valve to close slightly later, as venous return to the right side of the heart is increased. Inspiratory splitting of the second heart sound is the result – a normal finding in children and most adults. It is best appreciated with the stethoscope diaphragm applied at the left of the sternum in the second or third intercostal space.

Fixed splitting of the second heart sound is virtually pathognomonic of atrial septal defect. The second sound is split because of the increased volume load on the right ventricle, and the split is fixed because the septal defect equals right and left atrial pressure throughout the cardiac cycle. Fixed splitting is not a feature of ventricular septal defect or persistent ductus arteriosus.

Fixed splitting has to be distinguished from wide splitting of the second sound, where the split is audible in both inspiration and expiration but wider in inspiration. This occurs in right bundle-branch block and in pulmonary stenosis. There is a direct relationship between the width of the expiratory split and the pulmonary gradient. In severe pulmonary stenosis or Fallot's tetralogy, the pulmonary component of the second sound may be so quiet as to be inaudible. Wide splitting of the second sound is not usually found in pulmonary hypertension.

Reversed splitting of the second sound occurs when left ventricular ejection is prolonged or delayed so that pulmonary valve closure precedes aortic valve closure. Inspiration now causes the sounds to move together, so wider splitting is heard in expiration. In practice, reversed splitting is uncommon and is mainly associated with left bundle-branch block, hypertrophic obstructive cardio-myopathy and some cases of congenital aortic stenosis. In cases of severe aortic stenosis, the valve cusps are so rigid that the aortic second sound is diminished.

An abnormally loud second heart sound is most commonly due to systemic hypertension. The second sound may also be loud in patients with a dilated or aneurysmal ascending aorta. Because the pulmonary artery lies closer to the surface than the ascending aorta, pulmonary hypertension can cause a very loud pulmonary component of the second heart sound (P2), which may be palpable as well as audible, often accompanied by a right ventricular heave. In patients with transposition of the aorta and pulmonary arteries, a loud second sound is heard for the same reason.

The characteristics of the second heart sound in various conditions can be summarized as follows:

- Loud
- Thin chest wall
- Hyperdynamic circulation
- Pulmonary hypertension
- Transposition of the great arteries
- Soft
- Obesity
- Low cardiac output
- Severe aortic or pulmonary stenosis
- Normal split
- Healthy children, some adults
- Wide split
- Pulmonary stenosis
- Right bundle-branch block
- Fixed split
- Atrial septal defect
- Reversed split
- Hypertrophic cardiomyopathy
- Left bundle-branch block

THIRD SOUND

The third heart sound is a low-pitched sound, like a thump or a thud, which occurs in mid-diastole. It is 'physiological' in athletes, in some children, and in association with a hyperdynamic circulation, for example during pregnancy. In other patients, it is associated with a dilated, poorly contracting left ventricle with a high end-diastolic pressure. The precise
mechanism of the third heart sound is controversial. Its timing has been shown to coincide with the end of the phase of rapid diastolic ventricular filling and it is likely to be due to rapid acceleration and then deceleration of blood flow during early ventricular filling.

A third sound in a fit patient with a resting bradycardia is nearly always physiological. Likewise, when there is evidence of a hyperdynamic circulation such as loud first and second heart sounds, peripheral vasodilatation and a good pulse volume, a third sound is little cause for worry. A pathological third heart sound is heard in conditions characterized by high left or right ventricular end diastolic pressures. Common examples are heart failure (due to either preserved or reduced ejection fraction) and severe mitral regurgitation, even in the absence of heart failure.

A pathological third sound is usually part of a characteristic cadence described as a gallop rhythm. There is a tachycardia, a soft first heart sound quickly being followed by a soft second sound, and then a loud third sound: da-da-dum, da-da-dum. The patient often looks ill, and the cardiac apex is displaced and has a diffuse or dyskinetic feel. A chest radiograph will confirm cardiac enlargement, and the best way to confirm impaired ventricular function is by echocardiography.

FOURTH SOUND
The fourth heart sound is associated with atrial emptying into a stiff left ventricle. It is only heard in sinus rhythm, and precedes the first sound by about 0.15 seconds: da-lub-dup, da-lub-dup. It is characteristically heard in conditions that produce left ventricular hypertrophy and increased LV stiffness (e.g. hypertension and hypertrophic cardiomyopathy) and occasionally in patients with ischaemic heart disease, especially soon after myocardial infarction or aortic stenosis. In severe hypertension or hypertrophic cardiomyopathy, there is often a separate palpable and visible component to the apex beat that coincides with the fourth heart sound. A fourth sound is not a feature of mitral stenosis (where the stenosed valve prevents rapid atrial emptying) or of mitral regurgitation (where the atrium is too distended to contract forcefully). Some authorities describe patients with a fourth sound as having a ‘presystolic gallop rhythm’.

ADDED OR EXTRA SOUNDS
The opening snap is a feature of mitral (more rarely tricuspid) stenosis. It coincides with the bulging of the mitral valve ‘diaphragm’ into the ventricle in early diastole. It is a sharp, high-pitched sound best heard with the stethoscope bell in the third or fourth left intercostals space about 3 cm from the left sternal edge, and it is apt to be confused with a widely split second sound. The presence of an opening snap indicates that the valve, although stenosed, is still mobile. The interval between the second sound and opening snap reflects left atrial pressure – a high pressure, and thus severe stenosis, causes an early opening snap.

An ejection click is a short, loud, ringing sound immediately following the first heart sound (‘as the I follows c in click’). It is a feature of valvar aortic stenosis with a mobile aortic valve, bicuspid aortic valve, or valvar pulmonary stenosis, and it is usually followed by an ejection systolic murmur. It is best heard in the aortic or pulmonary ‘areas’ depending on its cause. The mechanism is thought to be tensing of the aortic or pulmonary cusps just prior to ejection. An ejection click without a murmur sometimes occurs in idiopathic dilatation of the pulmonary artery.

Mid-systolic clicks are usually associated with mitral valve prolapse (see HEART, MURMURS IN, p. 269) and are due to sudden tensing of parts of the mitral valve apparatus during systole. There may or may not be an associated systolic murmur. Both clicks and murmur may vary with posture and phase of respiration.

A clicking pneumothorax occurs when a small left pneumothorax causes a clicking sound, often loud and audible to the patient, in phase with the cardiac cycle. It is benign and self-limiting but may recur.

Prosthetic valve sounds are heard in patients who have undergone valve replacement with mechanical prostheses (e.g. Starr–Edwards, Bjork–Shiley or St Jude valves). Each valve has an opening sound (analogous to the opening snap or ejection click) and a closing sound (analogous to the first or second heart sound). The closing sound is usually much the louder – if it accompanies the first sound, the patient has had a mitral valve replacement, and conversely for aortic valve replacement. The sound and cadence of the clicks are fairly constant for an individual patient, and sudden muffling of one or other prosthetic sound usually indicates prosthetic malfunction, perhaps due to thrombosis.

HEARTBURN
Simon Anderson
Retrosternal burning, rising from the epigastrium towards the throat, lies within the spectrum of dyspeptic symptoms. It is commonly ascribed to the reflux of acid and bile into the lower oesophagus secondary to inappropriate relaxation of the lower
Simple antacids or motility agents alone. Alginates for reflux, being superior to H2-receptor antagonists, Proton-pump inhibitors offer the best initial therapy present.

or weight loss), must undergo an urgent endoscopy to with alarm features (dysphagia, odynophagia, anaemia recent-onset reflux-like symptoms, particularly those nowadays. Patients over the age of 55 years with (Bernstein test) – although this is rarely performed administration of dilute acid into the distal oesophagus (heartburn with normal acid/bile reflux into the distal oesophagus). It is found in approximately 5 per cent of patients complaining of reflux and is equally as likely to be found in those with epigastric pain alone. It is important as 0.3 per cent of people with Barrett’s oesophagus develop adenocarcinoma per year; those with high-grade dysplasia on biopsy have a foci of intramucosal cancer in 50 per cent of cases.

Acid reflux has also been implicated in the marked rise in the incidence of adenocarcinoma of the gastro-oesophageal junction reported from developed countries. The majority of people with significant acid reflux, however, do not have any symptoms; those with heartburn are no more likely than those with epigastric pain alone. It is important as 0.3 per cent of people with Barrett’s oesophagus develop adenocarcinoma per year; those with high-grade dysplasia on biopsy have a foci of intramucosal cancer in 50 per cent of cases.

HEMIANOPIA

Reginald Daniel

Hemianopia means an inability to see objects in one-half of the visual field. An understanding of the different types of hemianopia requires an understanding of the anatomy of the visual pathways (Fig. H.12). It should be noted that that the nerve fibres from the nasal half of each retina cross over at the optic chiasma, but those from the temporal sides do not. Thus, fibres from the nasal half of the left eye and the temporal half of the right eye form the right optic tract; and the fibres from the nasal half of the right eye and the temporal half of the left eye form the left optic tract.

MONOCULAR HEMIANOPIA

The loss of half of the visual field in one eye usually indicates optic nerve pathology, but may occasionally result from retinal damage. Monocular hemianopia may be either temporal or nasal, depending on which fibres in the optic nerve have been damaged. A monocular quadrantanopia indicates loss of one-quarter of one field.

Compression of the optic nerve from a tumour may result in a monocular hemianopia, but optic neuritis – such as frequently occurs in multiple sclerosis – more commonly produces a central field defect in the form of a scotoma. The most common cause of monocular hemianopia is ischaemic damage to the optic nerve.
BITEMPORAL HEMIANOPIA

This most commonly results from damage to the optic chiasma, usually from a pituitary tumour. Less common causes are suprasellar cysts, aneurysms, meningiomas of the tuberculum sellae or craniopharyngiomas.

When the pressure upon the chiasma begins from below, the visual field defect typically begins in the upper part of the temporal fields, leading initially to a bitemporal quadrantanopia. Pressure from above the chiasma initially produces a lower bitemporal defect. By the time all the crossing fibres in the optic chiasma are compressed, a complete bitemporal hemianopia will have developed.

Skull X-rays will usually show an enlarged pituitary fossa when a bitemporal hemianopia has resulted from a pituitary tumour. When bitemporal hemianopia is due to other causes, the skull X-ray may be normal. The investigation of choice is computed tomography scanning or magnetic resonance imaging (Fig. H.13).

HOMONYMOUS HEMIANOPIA

In homonymous hemianopia, the visual loss affects the right or left half of each visual field, usually resulting from damage to the optic radiation or visual cortex. Optic tract damage or damage to the lateral geniculate body may produce a homonymous hemianopia, but such cases are rare. Homonymous hemianopia may be either congruous or incongruous. Congruous defects are those which are identical in shape and degree in each field; incongruous defects are asymmetrical.

In the optic tracts, the intermingling of fibres from the homonymous halves of the two retinas is not as complete as in the optic radiation. Consequently, a lesion of the tract tends to lead to an incongruous homonymous hemianopia, whereas a lesion of the optic radiation is more likely to produce a congruous field defect.

Damage to the visual cortex may spare the area responsible for the macular vision, resulting in homonymous hemianopia with macular sparing. Lesions of the parietal lobe typically produce a hemianopia that begins with loss of vision in the lower parts of the field, whereas a temporal lobe lesion characteristically affects the upper parts of the visual field (homonymous quadrantanopias). Associated signs may assist in localization.

The visual field defect, when plotted using the Bjerrum screen, may be the same for different-sized objects, and it is then said to have ‘sharp edges’. A visual field defect that varies depending on the size of the object presented is said to have ‘sloping edges’. The former is more typical of a vascular cause, whereas the latter
is said to be more often encountered when the optic pathway is compressed by tumour.

In testing the visual fields clinically, it is customary to present an object – usually a finger or coloured pinhead – first to the temporal field of one eye, and then to that of the other eye. It is also useful to present a wagging finger to both temporal fields simultaneously. Occasionally, there is a failure in this test to pick up the finger in one field, although it is perceived when presented singly. This is the phenomenon of visual inattention (or extinction), and is encountered in some parietal lobe lesions.

HEMIPLEGIA

Mark Kinirons

Hemiplegia refers to complete paralysis of one side of the body affecting both the upper and lower limbs; hemiparesis refers to partial paralysis, although in practice the terms tend to be used interchangeably. The terms are usually applied in cases in which the paralysis is of upper motor neurone type, although it is possible to see unilateral weakness affecting both the arm and leg in lower motor neurone disorders, such as poliomyelitis, motor neurone disease or combined cervical and lumbar radiculopathy. Hemiplegia is most commonly seen in damage to the upper motor neurone above the level of the foramen magnum. A discrete lesion of the spinal cord in the upper cervical region may produce a hemiplegia, but this is rare.

Hemiplegia occurs most commonly as a result of ischaemic (Fig. H.14) or haemorrhagic (Fig. H.15) stroke affecting the motor cortex, internal capsule or, occasionally, the brainstem. The widely used Oxfordshire Community Stroke Project classification of stroke, which is predominantly clinical but refined by neuroimaging, is shown in Box H.17. Intracranial mass lesions such as tumour or subdural haematoma (Fig. H.16) may also present with hemiplegia, although usually of gradual onset. Typically, the so-called ‘physiological’ flexors are weaker than the extensors. This results in more profound weakness of shoulder abduction, elbow extension, wrist extension, finger extension and finger abduction in the arm, and hip flexion, knee flexion and ankle dorsiflexion in the leg. The opposing muscles tend to be those which are associated with spasticity, and this results in the typical hemiplegic posture of an abducted flexed arm and an extended leg. Associated signs include hyperreflexia, extensor plantar responses and ankle and knee clonus. The abdominal and cremasteric reflexes are lost on the side of the hemiplegia. In cases of chronic hemiplegia,
there may be some slight loss of muscle bulk, and contractures may become prominent. Paralysis occurring in childhood before bone growth ceases may be associated with hemi-atrophy.

The pathological causes of hemiplegia are listed in Table H.3.

### Table H.3 The causes of hemiplegia

<table>
<thead>
<tr>
<th>Category</th>
<th>Cause</th>
</tr>
</thead>
<tbody>
<tr>
<td>Congenital</td>
<td>Porencephaly, cerebral agenesis, cerebralangioma, Sturge–Weber syndrome (cerebral palsy)</td>
</tr>
<tr>
<td>Head injury</td>
<td>Birth injury, cerebral contusion, traumatic cerebral haemorrhage, subdural haematoma, extradural haematoma</td>
</tr>
<tr>
<td>Cerebrovascular accidents</td>
<td>Cerebral infarction and cerebral haemorrhage</td>
</tr>
<tr>
<td>Neoplasms</td>
<td>Primary neoplasms, secondary neoplasms</td>
</tr>
<tr>
<td>Infection</td>
<td>Meningitis (various), cerebral abscess, cortical thrombophlebitis, encephalitis, hydatid cyst</td>
</tr>
<tr>
<td>Demyelinating conditions</td>
<td>Multiple sclerosis, Schilder’s disease, acute disseminated encephalomyelitis</td>
</tr>
<tr>
<td>Degenerative conditions</td>
<td>Motor neurone disease</td>
</tr>
<tr>
<td>Miscellaneous conditions</td>
<td>Pick’s disease, epiloe</td>
</tr>
<tr>
<td>Functional</td>
<td></td>
</tr>
</tbody>
</table>

HICCOUGH

Alex West

Hiccough is very common and is only significant if persistent. It is caused by sudden involuntary contraction of both the diaphragm and the external intercostal muscles, in association with rapid glottic closure. Hiccoughs may be induced by stimulating a variety of sensory nerves, particularly the vagus and phrenic. They usually occur with a frequency of between 4 and 60 per minute.

The most common cause of hiccough is gastric distension after the rapid ingestion of food, alcohol or air. Other common causes include excitement and a sudden change in temperature, either of the environment or of the stomach, induced by a hot or very cold meal. However, in persistent cases, the following causes should be considered.

### INTRATHORACIC

#### Mediastinal

Irritation of a phrenic nerve may cause recurrent attacks of intractable hiccoughs. In an adult, this may
be due to malignant lymphadenopathy resulting from tumours of the lung or oesophagus, or to a lymphoma. An aortic aneurysm may rarely cause hiccough. Hiccough may also follow mediastinal surgery or mediastinitis.

**Diaphragmatic**
Hiccough may occur in pneumonia and empyema due to diaphragmatic pleurisy, and may also occur in myocardial infarction and pericarditis.

**Intra-abdominal disease**
Hiccough of intra-abdominal cause often results from diaphragmatic irritation due, for example, to diaphragmatic hernia, subphrenic abscess, peritonitis, pancreatitis, liver metastases, liver abscess, splenic infarct and carcinoma of the stomach. However, other conditions, such as carcinoma of the sigmoid colon and carcinoma of the uterus, have been associated with hiccough even without any obvious diaphragmatic involvement. Hiccough may occur after abdominal or pelvic operations and is seen in association with acute postoperative dilatation of the stomach and with intestinal obstruction.

**CENTRAL**

**Epidemic encephalitis**
This rare cause of hiccough was probably a variety of encephalitis lethargica in which there was inflammation affecting the third and fourth segments of the cervical cord, those from which the phrenic nerve originates. Hiccough may also follow lesions of the medulla.

**Intracranial**
Hiccough may rarely result from intracranial tumours, intracranial haemorrhage, brain abscess or meningitis, especially when the brainstem or basal meninges are involved.

**Toxic**
Uræmic hiccoughs are uncommon but may be persistent. Acute severe fevers, including typhoid, malaria and cholera, may also be accompanied by hiccoughs.

**Hysterical hiccough**
This occasional cause of hiccough usually leads to hiccough during wakefulness. However, the hiccough stops during sleep.

**Drugs**
Benzodiazepines and short-acting barbiturates can also cause hiccough.

**HIRSUTISM**
Paul Carroll
Hirsutism is said to be present when a female has an excessive growth of hair in androgen-sensitive areas where hair growth is normally minimal or absent. Hirsutism in isolation (i.e. without other signs of virilization) is usually benign, and the majority of patients are suffering from polycystic ovary syndrome (PCOS) or genetic influences. However, it is worth while looking for evidence of virilization (clitoral enlargement, deepening of the voice, temporal balding, decreased breast size and loss of the female body contour), which makes a pathological cause much more likely. The cause of hirsutism is an increased secretion of androgens from the ovary or adrenal glands. Normally, testosterone is produced directly from the ovary, and by extraglandular conversion of androstenedione secreted from both the ovary and the zona reticularis of the suprarenal cortex. The adrenal gland produces only minimal amounts of testosterone, but it is the main source of the pro-androgen dehydroepiandrosterone (DHEA) and its sulphate (DHEAS). Serum testosterone is markedly elevated in ovarian causes of virilization, and DHEAS is elevated in cases of virilization due to suprarenal pathology.

**TYPES OF HAIR GROWTH**

**Lanugo**
This is the very fine, silky, but sometimes quite long hair that covers the entire skin surface of the fetus; it is usually shed by the seventh or eighth month of intrauterine life.

**Vellus**
Males and females are born with the same number of hair follicles. In childhood, the follicles produce vellus hair, which is fine and non-pigmented. This type of hair persists into adult life in the areas of skin that are not producing terminal hair (see below). Occasionally, a woman may notice vellus hair, particularly on her face, when looking into a mirror with the sun or light behind her head. She may then look for them elsewhere and be surprised to find them on her chest, arms and legs. This, of course, is not hirsutism but 'pseudohirsutism'. Non-androgenic causes of hirsutism usually result in increased vellus hair formation.

**Intermediate hair**
This hair is soft and silky, but it may grow long and become pigmented. It then becomes a source...
of embarrassment. A combination of vellus and intermediate hair over the face and shoulders is characteristic of Cushing’s syndrome.

Terminal hair
Terminal hair is coarse and pigmented and is of three types:

- **Non-sexual hair**, which is present on the scalp, eyebrows, arms and legs
- **Ambosexual hair**, which is initiated by low levels of androgens and is present in the axillae, in the lower pubic triangle and on the limbs
- **Sexual hair**, which is produced by male levels of androgens, and is present in the upper pubic triangle as the ‘male escutcheon’, and on the face, nose, ears, trunk and limbs, where it is present in more profusion than ambosexual hair. When this type of sexual hair is present in a woman, she is deemed to be suffering from ‘hirsutism’

However, it is important to realize that there is an overlap between what can be considered to be normal and that which is regarded as ‘hirsute’. In a study of 400 consecutive Welsh and English women students at the University of Wales, 26 per cent had terminal hair on the face, 17 per cent on the chest or breasts, 35 per cent on the lower abdomen (mainly the linea alba), and 84 per cent on the lower arm and leg. Of these latter 84 per cent, nearly three-quarters also had terminal hair on the thighs and upper arms. As mentioned before, Mediterranean (and some Indian) women tend to grow more terminal hair than Nordic women, whereas women of the Mongolian races (Japanese, Chinese, American Indians, etc.) grow much less. These racial factors must therefore be taken into account when assessing hirsutism. The presence of oligomenorrhea, amenorrhea, and seborrhea and acne in a hirsute woman represents factors indicating the necessity for investigation.

**HIRSUTISM AND VIRILIZATION**
Virilism may accompany hirsutism and is characterized by clitoral enlargement, breast atrophy, temporal hair recession, frontal baldness and loss of the normal female contours due to increased muscularity. Deepening of the voice often occurs, and there is usually amenorrhea. The presence of virilization should draw attention to the possibility of one of the conditions marked by an asterisk in Box H.18.

**POLYCYSTIC OVARY SYNDROME**
PCOS is the most common cause of hirsutism in clinical practice. The classic features originally described by Stein and Leventhal are hirsutism, obesity, oligomenorrhea or amenorrhea, and enlarged cystic ovaries with thickened capsules. On microscopic examination, numerous small atretic follicles are found, surrounded by hyperplastic theca interna. In most cases, the menstrual disturbance starts shortly after puberty and tends to get progressively worse. Anovulation is invariable. Hirsutism is present in about 50 per cent of cases, and virilization is occasionally seen. The ovary produces excess androstenedione, which is converted into testosterone. In addition, the androstenedione undergoes conversion to oestrone in fatty tissue. Thus, there is not only androgenization but also continuous oestrogen production. The result is continuous high luteinizing hormone production by the pituitary, which tends to perpetuate the situation. Follicle-stimulating hormone levels are low normal. 17-Oxo- or ketosteroid levels are usually in the high-normal range.

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**Box H.18 Causes of hirsutism (in order of frequency)**

<table>
<thead>
<tr>
<th>Category</th>
<th>Causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Drugs</td>
<td>Menopausal preparations (containing methyltestosterone)</td>
</tr>
<tr>
<td></td>
<td>Androgens*</td>
</tr>
<tr>
<td></td>
<td>Anabolic steroids*</td>
</tr>
<tr>
<td></td>
<td>Synthetic 17-nor-progestogens; adrenocorticotropic hormone</td>
</tr>
<tr>
<td></td>
<td>Phenyltoxin†</td>
</tr>
<tr>
<td></td>
<td>Diazoxide†</td>
</tr>
<tr>
<td></td>
<td>Minoxidil†</td>
</tr>
<tr>
<td></td>
<td>Glucocorticoids†</td>
</tr>
<tr>
<td>Metabolic disorders</td>
<td>Anorexia nervosa†</td>
</tr>
<tr>
<td></td>
<td>Porphyria cutanea tarda†</td>
</tr>
<tr>
<td>Miscellaneous disorders</td>
<td>Cornelia de Lange syndrome†</td>
</tr>
<tr>
<td></td>
<td>Hypertrichosis lanuginosa†</td>
</tr>
<tr>
<td>Endocrine disorders</td>
<td>Acromegaly</td>
</tr>
<tr>
<td></td>
<td>Cushing’s syndrome*</td>
</tr>
<tr>
<td></td>
<td>Congenital and juvenile hypothyroidism*</td>
</tr>
<tr>
<td></td>
<td>Congenital adrenal hyperplasia*</td>
</tr>
<tr>
<td></td>
<td>Adrenal carcinoma or adenoma*</td>
</tr>
<tr>
<td></td>
<td>Ovarian tumours*</td>
</tr>
<tr>
<td></td>
<td>Arrhenoblastoma</td>
</tr>
<tr>
<td></td>
<td>Hilus cell tumour (and hyperplasia)</td>
</tr>
<tr>
<td></td>
<td>Luteoma</td>
</tr>
</tbody>
</table>

*May be associated also with virilization
†Associated with vellus rather than with terminal hair production
The classic syndrome may not always be present. Obesity may be absent, hirsutism is not a feature in 50 per cent of patients, and some patients may have normal-sized ovaries with a solitary atretic follicle. In contrast, the triad of hyperandrogenism, insulin resistance and acanthosis nigricans forms a specific subset of women with PCOS (HAIR-AN syndrome). In these women, the hirsutism is severe.

**IDIOPATHIC HIRSUTISM**

This is the second most common type of hirsutism, and it may reflect an increased sensitivity of the pilosebaceous unit to relatively normal plasma levels of androgen. The diagnosis should only be reached after excluding all the other possible causes listed above. If there is a history of hirsutism in other close female relatives, or a history of baldness in the father or brothers, the diagnosis of idiopathic hirsutism is more certain. Although the condition is called ‘idiopathic’, minor hormonal abnormalities are frequently found. In healthy women, small amounts of the pro-androgens androstenedione and DHEA are secreted mainly by the adrenal gland. These are then converted into testosterone and its derivative, the more powerful hormone dihydrotestosterone. In women with hirsutism, there may be an increased production of androstenedione by the ovary as well as the adrenal, and this leads to increased conversion to testosterone and dihydrotestosterone. The rise in active androgens lowers the concentration of sex hormone-binding globulin (SHBG); the net effect is to decrease the amount of bound androgen and to increase the amount of free androgen, thus amplifying its biological action. In idiopathic hirsutism, the usual finding is either a slightly raised or a normal plasma testosterone. If the SHBG is reduced, the free testosterone may be elevated. Urinary 17-keto- or oxosteroid levels are usually normal, because they are increased only when very large amounts of androstenedione and DHEA are being produced.

**MENOPAUSE**

At the time of the menopause, there tends to be an increase in the growth of terminal hair in the moustache and beard areas, whereas, paradoxically, body hair gradually becomes less. This is physiological because of the change in the oestrogen:androgen ratio.

**DRUGS**

Menopausal preparations containing methyltestosterone, androgens in general, anabolic steroids, the synthetic 17-nor-progestogens and adrenocorticotropic hormone (ACTH) may all cause an increased growth of terminal hair. Androgens and anabolic steroids may even cause some degree of virilization.

Other drugs (phenytoin, diazoxide, minoxidil and glucocorticoids) cause stimulation of vellus hair growth.

**METABOLIC AND MISCELLANEOUS DISORDERS**

Vellus hair growth is seen as a feature of anorexia nervosa and porphyria cutanea tarda. In the condition known as hypertrichosis lanuginosa, the lanugo hair that was shed _in utero_ at the seventh or eighth month suddenly regrows and completely covers the face, so that the features become unrecognizable. In Cornelia de Lange syndrome (see STATURE, SHORT p. 645), which is associated with short stature ('Amsterdam dwarfism'), the scalp hair reaches down to the bushy and confluent eyebrows, and there is also generalized hirsutism of a mixed vellus and intermediate type.

**ENDOCRINE DISORDERS**

**Acromegaly**

The incidence of this disease is approximately six newly diagnosed cases per million of the population each year, whereas the prevalence of previously diagnosed cases is approximately 40 per million of the population. Just over half the women with acromegaly complain of hirsutism, and in them urinary 17-oxosteroids may be slightly raised. The mechanism for the increase in 17-oxosteroids is unknown. The hair is of the terminal type.

**Cushing’s syndrome**

Apart from iatrogenic disease caused by the administration of glucocorticoids, Cushing’s syndrome is rare, occurring in only one or two per million of the population each year. The condition is relatively more common in women, when it is usually associated with hirsutism. There is conspicuous hair growth in the beard and moustache areas, but the characteristic feature is a rather widespread, vellus hirsutism on the face and over the shoulders. The plethoric moon-shaped face, buffalo hump, supraclavicular puffiness, livid cutaneous striae, central obesity, thin limbs and subcutaneous bruising of the classic case make the diagnosis easy. However, mild or early cases of Cushing’s syndrome may not be easy to recognize, and it is a diagnosis always worth bearing in mind in a woman with hirsutism. The best screening tests to use are a 24-hour urine collection for free cortisol or a 9 a.m. plasma cortisol following 1 mg of oral dexamethasone taken the previous night before retiring. A normal urinary free cortisol and a suppressed 9 a.m. plasma cortisol will exclude the diagnosis.

**Congenital and juvenile hypothyroidism**

Congenital hypothyroidism occurs in approximately 1 per 5000 births in the UK. With the increasing use...
of neonatal thyroid-stimulating hormone screening, most cases should be diagnosed shortly after birth and treatment initiated. In cases diagnosed later, the babies may develop a coat of vellus or intermediate hair over the back and extensor surfaces of the skin. A similar picture is sometimes seen in patients with juvenile hypothyroidism.

**Congenital adrenal hyperplasia**

Congenital adrenal hyperplasia (CAH) results from excessive ACTH stimulation. This is caused by an inherited deficiency of an enzyme necessary for the synthesis of cortisol. Five different types of enzyme lack have been recognized, but 21-hydroxylase deficiency is by far the most common. This occurs with a frequency of 1 per 5000 live births. In the female, the condition is usually recognized at birth because of genital abnormalities such as enlargement of the clitoris and fusion of the labia. In girls with less severe enzyme defects, 21-hydroxylase may present at a later age with premature false puberty (see PUBERTY, PRECOCIOUS p. 536) and hirsutism. Occasionally, the condition becomes apparent after puberty; in addition to hirsutism, there may be clitoromegaly and either oligomenorrhea or amenorrhea.

The increase in adrenal androgens is caused by an ACTH-stimulated build-up of precursors of cortisol. They can be measured as 17-oxosteroids in the urine. In 21-hydroxylase defect, raised levels of 17-hydroxy-progesterone are found in the blood, and its metabolite, pregnanetriol, is increased in the urine. Women with less severe forms of CAH present later than the childhood period, typically in adolescence when puberty is completing. These women are considered to have the non-classic form of congenital adrenal hyperplasia, and the diagnosis can be confirmed by the demonstration of a marked rise in 17-hydroxyprogesterone following ACTH administration.

**Adrenal carcinoma or adenoma**

Adrenal tumours producing hirsutism and virilization are rare, and symptoms and signs develop abruptly. There are usually very high levels of urinary 17-oxosteroids that cannot be suppressed to 50 per cent of the original levels by 0.5 mg of dexamethasone 6-hourly for 3 days. Forty per cent of carcinomas are palpable. Adenomas may be very small. Calcification in a carcinoma may sometimes be seen on X-ray. Computed tomography scanning is usually informative but, if not, isotope scanning or selective venous sampling for adrenal androgens may localize the tumour.

**Ovarian tumours**

These are exceedingly rare as a cause of hirsutism and virilism, and symptoms and signs develop abruptly. They usually give rise to extreme degrees of virilism because the androgen that they produce, testosterone, is very potent. Levels of testosterone are high in the blood, but urinary 17-oxosteroids, which only measure 25 per cent of all testosterone secreted, may be within the normal range. The tumour may be palpable on pelvic examination and the enlargement can be confirmed by ultrasound or magnetic resonance imaging.

### HYPERVENTILATION

Andrew Hodgkiss

Hyperventilation is overbreathing or ventilation in excess of metabolic requirements, with symptoms arising from respiratory alkalosis and hypocapnia (Box H.19).

**ACUTE HYPERVENTILATION**

This condition is easily recognized because overbreathing is apparent. The characteristic patient is a young woman, and the presentation has a readily identified precipitant that may have aroused anxiety, fear, distress, excitement, adulation or other powerful feelings. Occasionally, the respiratory alkalosis is compensation for a metabolic acidosis, and especially in the young, diabetic ketoacidosis and salicylate poisoning can present in an occult manner and ought to be excluded. Asthma, which has been identified as both a cause and a consequence of hyperventilation, may also present as stress-induced

### Box H.19 Causes of acute hyperventilation

<table>
<thead>
<tr>
<th>Most common</th>
<th>Rare</th>
</tr>
</thead>
<tbody>
<tr>
<td>- Stress reaction</td>
<td>- Acidotic disorders</td>
</tr>
<tr>
<td>- Anxiety</td>
<td>- Biguanide intolerance</td>
</tr>
<tr>
<td>- Phobia</td>
<td>- Intestinal fistula</td>
</tr>
<tr>
<td>- Panic</td>
<td>- Surgical relocation of ureters in ileum or colon</td>
</tr>
<tr>
<td><strong>Less common</strong></td>
<td><strong>Others</strong></td>
</tr>
<tr>
<td>- Acidotic disorders</td>
<td>- Epilepsy</td>
</tr>
<tr>
<td>- Diabetic ketoacidosis</td>
<td>- Acute porphyria</td>
</tr>
<tr>
<td>- Salicylate poisoning</td>
<td>- Phaeochromocytoma</td>
</tr>
<tr>
<td>- Renal failure</td>
<td>- Hepatic cirrhosis</td>
</tr>
<tr>
<td>- Pulmonary disorders</td>
<td>- Pyloric stenosis</td>
</tr>
<tr>
<td>- Asthma</td>
<td></td>
</tr>
<tr>
<td>- Pulmonary oedema</td>
<td></td>
</tr>
<tr>
<td>- Pulmonary embolism</td>
<td></td>
</tr>
<tr>
<td>- Pleural effusion</td>
<td></td>
</tr>
<tr>
<td>- Spontaneous pneumothorax</td>
<td></td>
</tr>
</tbody>
</table>
dyspnoea, although it is usually distinguishable by evident expiratory difficulty. In hospital, acute hyperventilation is not uncommonly precipitated by abdominal surgery or childbirth, when anxiety is accompanied by an avoidance of abdominal breathing as a protective mechanism against pain or fear of re-opening the wound. The differential diagnosis of acute breathlessness in these circumstances is pulmonary embolism, which may also provoke as well as mimic acute hyperventilation.

**CHRONIC HYPERVENTILATION**

Chronic hyperventilation syndrome is much more difficult to recognize because, once established, the disorder can be maintained by occasional deep breaths or sighs. Chronic hyperventilation accounts for 6 per cent of medical referrals, but it is frequently unconsidered, with the patient’s multiple unexplained somatic and psychological complaints leading all too readily to a label of ‘medically unexplained symptoms’. Even when chronic hyperventilation syndrome is contemplated, the clinician faces difficulty establishing the disorder’s presence because there is no satisfactorily reliable sign or test. The diagnosis rests upon a combination of typical symptoms and signs, clinical tests and investigations of respiratory function. The central symptom is breathlessness, an air hunger that is associated with gasping, sighing and a feeling of suffocating. Breathlessness tends to be worse at rest, following exercise or on awakening, and it may be associated with a particular setting, such as shops or buses, or be triggered by emotions. Common physical symptoms include fatigue, dizziness, faintness, headaches, tremors, sweating, palpitations, chest pain, dysphagia, nausea, heartburn, diarrhoea and flatulence. Common psychological symptoms are anxiety, depression, poor concentration and memory and depersonalization. Any of these may be the presenting complaint, but it is the sheer number and range of symptoms that provide the clue. A more characteristic pointer is paraesthesiae in the hands, fingers and around the mouth, while tetany is strong (but rare) evidence. In susceptible individuals, hyperventilation may present with fits.

At interview, gasping, sighing and a rapid, uneven respiratory pattern may be noted. The frequent interruption of speech by the need to breathe can be an important sign. Excessive thoracic movement during breathing is characteristic, while forced, voluntary overbreathing may rapidly provoke the re-experience of presenting symptoms; re-breathing into a paper bag may abolish these. Assessment of respiratory function helps to establish the diagnosis (Box H.20).

It cannot be assumed that chronic hyperventilation invariably arises from a disturbed emotional state. Physical causes include disorders of ventilation–perfusion: asthma, pulmonary embolism, parenchymatous lung disease and, rarely, neurological damage to the respiratory pathways, usually in the brainstem. However, the most common causes are psychological disorders, especially anxiety, phobias and panic disorder. Paradoxically, it is now evident that chronic overbreathing may induce these conditions as well as result from them. This interaction is complex and best understood as a feedback loop (Fig. H.17), although the implication is that treatment can be focused upon either the breathing abnormality or the emotional disorder with the expectation of ameliorating both aspects.

Finally, a substantial minority have no evidence of underlying physical or psychological disorder. In these patients, chronic hyperventilation syndrome is probably a primary habit disorder, although the predisposing biological and psychosocial mechanisms are still unknown. All habits tend to be exaggerated when under pressure, so it should be anticipated that such individuals become symptomatic on stressful

**Box H.20 Causes of chronic hyperventilation**

<table>
<thead>
<tr>
<th>Most common</th>
<th>Rare</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anxiety</td>
<td>Neurological disorders</td>
</tr>
</tbody>
</table>
| ![hyperventilation](image)

Figure H.17 The relationship between hyperventilation and anxiety.
HYPOCHONDRIASIS

Andrew Hodgkiss

The characteristics of hypochondriasis are: (i) a concern with health or disease in oneself, present most of the time; (ii) the preoccupation not being justified by the amount of organic pathology; and (iii) the individual not responding adequately to reassurance, given after a careful physical assessment. The abnormal health beliefs lie on a spectrum ranging from a preoccupation to an overvalued idea to a delusion. Hypochondriacal delusions have a different differential diagnosis and require different treatment from non-delusional hypochondriasis.

When faced with a hypochondriacal patient, the first and most essential step is to defer reaching this conclusion until physical conditions are excluded. Suspicions that the presentation is not hypochondriacal are raised if (i) the presentation is acute; (ii) there is no past history of medically unexplained symptoms; (iii) there are no stressors evident; and (iv) there is no psychological disturbance beyond the anxiety/apprehension accompanying the stressors evident.

Hypochondriacal presentations are usefully divided into primary and secondary, the distinction made more important by the poor prognosis found in primary hypochondriasis contrasting with the favourable outcome generally attained when hypochondriasis is in the setting of another psychological disorder.

Transient hypochondriacal reactions are common, usually responding well to explanation and reassurance, but occasionally developing and persisting. Any stress or situation may be the trigger in a vulnerable individual, but there are some common circumstances in which personal vulnerability is less crucial. Recovering from serious illness may precipitate this response as part of an adjustment reaction; similarly, serious illness in a close friend or relative may lead to concerns about having the same disease, the mental mechanism being identification with the loved one and sharing their suffering, or as part of a grief reaction when death has occurred. Hypochondriacal reactions are more common in the elderly, particularly if the patient has an established painful condition, and in this group the loss of work, outside interests and friends may conspire to focus attention and interest on the functioning of the ‘inner world’.

Hypochondriacal delusions are uncommon and may take easily recognizable, bizarre forms in schizophrenia, dementia and monosymptomatic hypochondriacal psychosis (a rare condition in which the delusions are the only psychopathology found). Among the more common presentations are beliefs of infestation, emitting odours, and changes in the size, shape or function of the organs. In psychotic depression, the delusions may be more subtle and difficult to detect, although detection is the more important because these phenomena are associated with a substantial risk of suicide. Generally, depressive hypochondriacal delusions are understandable in the context of other depressive changes – for instance, delusions of venereal disease may arise from depressive guilt and erectile dysfunction, while delusions of cancer can be the outcome of morbid, pessimistic thinking accompanied by constipation, impaired appetite and weight loss.

Hypochondriacal preoccupations, worries and fears occur commonly in the setting of neurotic disorders. In anxiety, agoraphobia, panic disorder and non-psychotic depression, any hypochondriacal features can usually be easily established as one element in a constellation of symptoms and signs that typify the underlying disorder. Physical symptoms may be the presenting feature of any of these conditions, and they may be perceived with enhanced alarm or dread by a patient who has an abnormal mood state. The presenting complaint is usually a typical feature of anxiety or depression, such as musculoskeletal pain, cardiovascular, gastrointestinal or neurological symptoms, which basically becomes magnified and misinterpreted.

**Box H.21  Causes of hypochondriasis**

<table>
<thead>
<tr>
<th>Most common</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Stress reaction</td>
</tr>
<tr>
<td>• Adjustment reaction (especially to illness)</td>
</tr>
<tr>
<td>• Anxiety</td>
</tr>
<tr>
<td>• Depression</td>
</tr>
<tr>
<td>• Early or otherwise undetected physical illness</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Less common</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Panic</td>
</tr>
<tr>
<td>• Phobia</td>
</tr>
<tr>
<td>• Obsessive–compulsive disorder</td>
</tr>
<tr>
<td>• Schizophrenia</td>
</tr>
<tr>
<td>• Primary hypochondriasis</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Rare</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Dementia</td>
</tr>
<tr>
<td>• Mental impairment</td>
</tr>
<tr>
<td>• Schizoaffective disorder</td>
</tr>
<tr>
<td>• Monosymptomatic hypochondriacal delusional state</td>
</tr>
<tr>
<td>• Post-traumatic stress disorder</td>
</tr>
<tr>
<td>• Hysteria</td>
</tr>
<tr>
<td>• Briquet’s syndrome (somatization disorder)</td>
</tr>
</tbody>
</table>
Primary hypochondriasis is rare, but its existence as a diagnostic entity is now widely accepted. It is essentially a diagnosis by exclusion of the causes already outlined, but the condition has some characteristic features: (i) first and foremost, persistent complaining; (ii) detailed accounts: ‘the organ recital’; (iii) matter-of-fact presentation; and (iv) less interest in the response than in the narrating. Such patients are usually obsessional and often anxiety-prone. These characteristics of style and personality aid in distinguishing primary hypochondriasis from somatization disorder (Briquet’s syndrome) where patients have persistent unexplained physical complaints, but tend to be inarticate, poor historians who give highly dramatic accounts of events but sketchy information about their symptoms, with the emphasis upon extraordinariness rather than detail.

Finally, it should be borne in mind that patients with primary hypochondriasis are not immune to physical illnesses, and are indeed vulnerable to mental disorders: exaggeration in hypochondriacal complaining may indicate a superimposed depressive illness or anxiety state. On the other hand, repeated specialist referrals and investigations reinforce this behaviour. Hence, the doctor has a delicate balance to strike over when to investigate and treat, and unfortunately this decision-making is frequently hindered by the antipathy and alienation that tends to develop in the doctor–patient relationship.

Latterly, there has been increasing optimism about the treatability of primary hypochondriasis, specifically with cognitive–behavioural psychotherapy. Elements of this treatment include the banning of reassurance, self-monitoring and the challenging of abnormal health beliefs.

**HYPOTHERMIA**

Mark Kinirons

Hypothermia is defined as a core temperature of below 35 °C (95.8 °F). Normally protected parts (e.g. the abdomen) are cold to the touch. When hypothermia is suspected, the deep rectal temperature should be measured with a low-reading thermometer. It is important to have a low threshold of suspicion in older people who will not feel the cold, and therefore neither complain of it nor make concerted attempts to rewarm themselves.

**Mild hypothermia** down to 32 °C (90 °F) may be characterized by confusion, slow responses, slurred husky speech, ataxia and involuntary movements resembling cerebellar incoordination. Skin pallor, shivering and some degree of tachycardia and mild hypertension are often noted.

**Moderate to severe hypothermia** is characterized by drowsiness, stupor or overt coma. Some patients may be comatose at 31 °C; others may still be conscious at 27 °C. The pupils are usually constricted and unresponsive to light (mimicking opium poisoning or pontine haemorrhage). Occasionally, fixed dilated pupils are noted in those with severe, protracted hypothermia and then may mimic closed head injury. The skin may show cyanotic or pink blotches, and sometimes purpuric lesions, skin haemorrhages or pressure lesions. Fluid shifts give the skin a typical puffy or oedematous consistency. At very low core temperatures, fine muscle tremors and muscle rigidity have replaced shivering. Tendon jerks, if still present, show slow contraction and relaxation phases. The knee jerk is the last to be lost. The plantar responses may be bilaterally extensor or absent; a unilateral extensor response usually indicates an underlying cerebrovascular catastrophe. Severe bradycardia or atrial fibrillation may be present. Tachycardia is inappropriate in hypothermia and suggests internal haemorrhage, for example from severe gastric erosions. Any intra-abdominal catastrophe may not be detectable until after rewarming. Coarse pulmonary crepitations indicate oedematous and often infected lungs, but respiration may be too poor for these features to be recognized. An electrocardiograph (ECG) is likely to show characteristic J waves (Fig. H.18).

Severely hypothermic patients lack easily detectable vital signs, with imperceptible peripheral pulses, inaudible heart sounds, unrecordable blood pressure, and slow, shallow respiration; the pupils may be widely dilated and unresponsive. An ECG will exclude death, but even ventricular fibrillation may reverse with rewarming. The diagnosis of death should not be made until rewarming has been achieved and vigorous resuscitation attempts made.

Many elderly hypothermic patients have complex physical signs, either because of complications of the hypothermia (e.g. pulmonary crepitations) or because the risk of developing hypothermia is greater with concomitant medical disease. It should be remembered that, while hypothermia is more common during cold weather in older people, the condition is by no means confined to the colder winter months.

The important causes of hypothermia are listed in Box H.22.
Hysteria was one of the two psychoneuroses that preoccupied Freud in the late nineteenth century, the other being obsessive–compulsive neurosis. Freud famously insisted that the aetiology of both conditions was sexual experiences in childhood. The hysterical, usually female, was traumatized by the experience once they were old enough to recognize the abuse of power involved, while the obsessional, usually male, was left with protracted guilt for enjoying aspects of the experience.

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**Box H.22** Causes of hypothermia

<table>
<thead>
<tr>
<th>Decreased heat production</th>
<th>Increased heat loss</th>
<th>Central thermoregulatory failure</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Inactivity and immobility (e.g. lying on the floor after a stroke or fractured femur)</td>
<td>• Outdoors cold/wet exposure or cold-water immersion</td>
<td>• Uraemia</td>
</tr>
<tr>
<td>• Hypoglycaemia</td>
<td>• Low ambient home temperatures due to poor heating and housing</td>
<td>• Cerebrovascular injury or cerebral trauma</td>
</tr>
<tr>
<td>• Impaired shivering capacity (elderly)</td>
<td>• Reduced awareness of cold due to dementia or delirium</td>
<td>• Severe chronic or debilitating disease (preterminal) (e.g. malignancy, hepatic cirrhosis)</td>
</tr>
<tr>
<td>• Myxoedema</td>
<td>• Ethanol</td>
<td>• Overwhelming infection (e.g. severe bronchopneumonia, overwhelming tuberculosis, falciparum malaria)</td>
</tr>
<tr>
<td>• Starvation and malnutrition</td>
<td>• Vasoconstriction failure (e.g. elderly survivors of previous accidental hypothermia)</td>
<td>• Severe congestive heart failure</td>
</tr>
<tr>
<td>• Hypopituitarism</td>
<td>• Malnutrition (e.g. kwashiorkor)</td>
<td>• Shock after multiple injury, major operation, severe coronary thrombosis or pulmonary embolism</td>
</tr>
<tr>
<td></td>
<td>• Erythroderma (e.g. generalized psoriasis, exfoliative dermatitis, Paget’s disease of bone)</td>
<td>• Collapse due to severe dehydration (e.g. severe diarrhoea, including cholera)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>• Pancreatitis</td>
</tr>
</tbody>
</table>

**Mechanism(s) uncertain**

- Drugs (including self-poisoning) (e.g. phenothizine tranquilizers, tricyclic antidepressants, barbiturates)
- Diabetic ketoacidosis
- Wernicke’s encephalopathy
- Carbon monoxide poisoning/anoxic brain injury (e.g. after cardiac arrest, organophosphate poisoning)
- Blood transfusion reaction

**Peripheral thermoregulatory failure**

- Spinal cord lesions, especially cervical
- Autonomic neuropathy (e.g. diabetes melitus, Parkinson’s disease)

**Figure H.18** ECG showing characteristic J waves in a 67-year-old male with myxoedema (rectal temperature 25 °C).
Freud incorporated several of Charcot’s clinical observations in this account of hysteria – suggestibility, belle indifférence and, most notably, the way in which the distribution of signs followed the patient’s lay understanding of anatomy rather than real anatomy (e.g. the glove and stocking distribution of sensory loss or the weakness of a whole upper limb). Freud, originaly and crucially, observed that the nature and distribution of hysterical somatic symptoms was meaning-laden, symbolic and potentially interpretable in terms of the patient’s autobiography. For example, facial pain in a young woman rejected by her fiancée was experienced ‘like a slap in the face’. Freud felt that such symptoms served to relieve the intense psychological conflict lying behind them – the primary gain was this resolution of psychological conflict through somatic expression.

In the twenty-first century, we discuss Freudian hysteria as conversion disorder. A diagnosis of conversion disorder still requires positive evidence of a psychological conflict underlying and explaining the medically unexplained somatic symptom (Box H.23). We also draw a distinction between conversion symptoms (which are common in primary care and among depressed patients) and conversion disorder proper (which requires the conversion symptoms to have been present and stable for months). Conversion disorder is confined to sensori-motor symptoms. Limb weakness, aphony, astasia–abasia (i.e. difficulty standing and walking) and hysterical blindness are typical presentations. Strictly speaking, chronic pain that is medically unexplained should be considered separately (see PAIN, PSYCHOGENIC, p. 489).

Conversion disorder is more commonly seen in women than men, and the age of onset is nearly always under 30 years of age. Indeed, the onset of apparent conversion symptoms in a patient over 40 years of age demands intensive medical investigation. Occult malignancy or neurological disorder is highly likely in the older patient.

The differential diagnosis of conversion disorder is considered in Box H.24. Neurological disorders, systemic disorders with central nervous system effects (such as systemic lupus erythematosus) and occult malignancy should all be ruled out by physical examination and investigation. Conversion symptoms may occur prominently as part of the clinical picture in both depressive illness and schizophrenia.

Perhaps the most famously difficult diagnostic distinctions are between conversion disorder, factitious disorder and malingering. The textbook view is that malingering is conscious deception of the doctor for real-world gain (such as avoidance of military service or obtaining money) while conversion disorder is an unconscious process, and the gain is predominantly the primary gain Freud described (i.e. the resolution of a psychological conflict through its conversion into a somatic symptom). But matters are not so clear-cut, and any doctor who has tried to give expert testimony will confirm how difficult it is to be sure on these matters. Patients with conversion disorder do enjoy secondary gains (such as attention and care from their family, being excused from work and possibly the offer of financial benefits), and this can shade into exactly the type of real-world gain at stake in malingering. In the end, the matter is often a judgement on the honesty of the patient, and some expert medical witnesses are now asking the court, rather than the doctor, to decide this.

The textbook difference between conversion disorder and factitious disorder is that, in the latter, there is deliberate conscious deception of healthcare professionals in order to win unnecessary medical interventions that the patient finds a rewarding form of care. Patients will not be able to articulate why they find, for example, repeated abdominal surgery rewarding. But their repeated hospital presentations suggest that they do. The current view is that diagnosing factitious disorder requires forensic evidence of deception (such as CCTV footage or laboratory findings). The association with factitious disorder by proxy means that a careful assessment of risk to others must be undertaken in such cases, and this is a highly specialized task.
Indigestion (dyspepsia) is a vague term that encompasses a constellation of symptoms, including upper abdominal discomfort or bloating, retrosternal or epigastric pain, nausea, loss of appetite and early satiety. Dyspepsia affects 20–40 per cent of Western populations, 25 per cent of whom seek medical attention. In only 25 per cent of cases is a cause found. Certain associated symptoms indicate a strong possibility of an underlying serious cause. For possible gastrointestinal bleeding, the ‘alarm’ features are listed below:

- ≥55 years with either new onset dyspepsia; persistent dyspepsia despite adequate treatment; history of peptic ulcers or cancer; NSAID or aspirin use
- Dysphagia
- Weight loss
- Vomiting
- Iron deficiency/anaemia
- Epigastric mass
- Suspicious barium meal.

Many conditions may give rise to dyspepsia. The most common are acid-related conditions, comprising ulceration or inflammation of the oesophagus, stomach or duodenum, and non-ulcer dyspepsia. The latter is a variant of irritable bowel syndrome (IBS). Other conditions to be considered are gallstones, gastroparesis (diabetes mellitus), viral or drug-induced gastropathy, infiltrating conditions (eosinophilic enteropathy, lymphoma and scleroderma), pyloric outlet obstruction, Crohn’s disease, giardiasis, gluten enteropathy, pancreatitis, achalasia and space-occupying lesions of the upper gastrointestinal tract (adenocarcinoma, squamous cell carcinoma, mucosa-associated lymphoid tissue lymphoma, metastatic cancer, Kaposi’s sarcoma, stromal cell tumours, submucosal cysts, ectopic pancreas). Systemic disease and its therapy, or pathology elsewhere in the gastrointestinal tract (such as colitis), may cause marked nausea. Many of the conditions mentioned are intimated at by the clinical setting or are only diagnosed following investigation.

The main focus of investigating dyspepsia is to exclude cancer. The most accurate test is an endoscopy (oesophago-gastro-duodenoscopy or gastroscopy (OGD)). Because dyspepsia is so common, it is not practical to perform an endoscopy on all patients with dyspepsia. Most countries have recommendations, based on the local incidence of *Helicobacter pylori* infection (see below) and cancer. Examples include the British Society of Gastroenterology (www.bsg.org.uk) and the American Gastroenterological Association (www.gastro.org) guidelines.

*Helicobacter pylori* is a Gram-negative anaerobe that colonizes the gastric antrum in a variable proportion of the population, being more common in the elderly and those from poorer socioeconomic groups. Its presence is associated with more than 90 per cent of duodenal ulcers, 60–70 per cent of gastric ulcers, and an increased risk of the development of gastric mucosa-associated lymphoid tissue lymphoma and distal gastric adenocarcinoma (Fig. I.1). Those patients infected are more susceptible to non-steroidal anti-inflammatory drug-induced gastric injury than the general population. The association between *H. pylori* and reflux oesophagitis or non-ulcer dyspepsia, importantly, is uncertain but probably weak.

The antigenic properties of *H. pylori* and its ability to split urea are used to diagnose its presence through serology, stool sampling, carbon-13- or carbon-14-labelled urea breath test, or gastric mucosal biopsy. Patients under the age of 55 years with dyspepsia but no alarm features should be tested for *H. pylori*. Most guidelines recommend either the stool antigen test or C13/C14 urease breath test. The serum antibody test is sensitive but not specific (in particular it detects previous, resolved infection) and is not

![Figure I.1](image-url) Acute pre-pyloric antral ulcer with some oozing of fresh blood.
INFERTILITY

 verifies the presence of a cancer.

H. pylori

Testing and treating of

H. pylori

failure to conceive within 12 months of commencing unprotected regular intercourse. After 1 year, 80–85 per cent of couples have managed to conceive, and a further 5–10 per cent of pregnancies will have been successful by the end of the 18 months. It is estimated that approximately 1 in 7 couples will have infertility and require investigation. Infertility can be divided into:

- Primary, when the woman has never been pregnant
- Secondary, when the woman has had a previous pregnancy, whatever the outcome

It is essential that both partners be investigated. Causes of infertility are shown in Box I.1. The incidence of the causes of infertility can be summarized as follows:

- Tubal problems: 20 per cent
- Anovulation: 25 per cent
- Other female factors, uterine or peritoneal factors, e.g. endometriosis: 10 per cent
- Male factors: 30 per cent
- Sexual problems: 5 per cent
- Unexplained: 25 per cent

In approximately 40% of cases, a disorder will be found in both partners, which explains why the numbers in the list above do not add up to 100 per cent.

It is always important to explore the frequency of sexual intercourse since the sperm lasts for approximately 72 hours, during which it can fertilize the ovum, which is fertilizable for approximately 24 hours following ovulation. Therefore, timing may be a significant factor for some couples. In many patients, the failure is due to a number of minor ‘infertility factors’, unimportant in themselves, but which in aggregate may result in an inability to conceive.

Both partners will need investigation and no assumptions should be made. In all cases, the woman should be screened for Chlamydia trachomatis and susceptibility to Rubella before embarking on further investigations and treatment. As a general rule, referrals for fertility investigations are usually after 12 months but an earlier consultation should be offered in women over the age of 36 years due to declining fertility, or if there is a known cause for infertility or predisposing factors. It should be emphasized to the women that a body mass index (BMI) >30 or <19 may cause her to take longer to conceive.

The NICE guidelines (NICE guideline 156: Fertility: Assessment and treatment for people with fertility problems (2013) www.nice.org.uk/guidance/cg156) recommend the following investigations:

- Semen analysis
- Assessment of ovulation
- Assessment of tubal damage and uterine abnormalities

Semen analysis

Spermatogenesis takes 74 days to occur, and consequently the sperm count may reflect the health of the individual when that process commenced.
A routine semen analysis will look at the following factors:

- **Volume:** >1.5 ml
- **pH:** 7.2
- **Count:** >15 million/ml
- **Motility:** >32 per cent forwardly progressive
- **Morphology:** >4 per cent normal forms

As the sperm count varies in each ejaculate, an initial count that is found to be low should be repeated at least once. The ejaculate should be fresh and reach the laboratory within 1–2 hours. Providing there are some spermatozoa in the ejaculate, it is not possible to say that a man is infertile no matter how low the semen count, as pregnancies seem to occur in the face of very low counts. However, it is considered that the male factor is significant if two or more semen analyses have one or more variables less than the 5th centile as defined by the WHO (2010) criteria above.

**Ovulation factors**

When a woman ovulates will be dependent on her menstrual cycle length and her age. In a normal menstrual cycle of 28 days, ovulation will occur at 14 days. The time from ovulation to menstruation is the luteal phase of her cycle and lasts 14 days. The follicular phase (i.e. how long it takes for the follicle/egg to develop) is the variable factor. Ovulation can be checked by a mid-luteal phase serum progesterone, which should be over 30 nmol/l. Basal body temperature charts are not helpful and should be avoided as they can increase the woman's stress. Ovulation kits are now available over the counter but can be expensive; they are used to test the urine for the luteinizing hormone (LH) surge.

In irregular cycles, the follicle-stimulating/luteinizing hormone (FSH/LH) levels should be checked. Prolactin should only be checked if there is an ovulatory disorder, galactorrhoea or suspicion of a pituitary tumour. Thyroid function tests should only be performed if there are symptoms suggestive of thyroid disease.

Age is a good predictor of ovulation as fertility reduces with age, but in order to get an idea of ovarian response to in-vitro-fertilisation techniques, one of the following investigations is recommended:

- A total antral follicle count of 4 or less suggests a poor response; more than 16 suggests a good response.
- Anti-Müllerian hormone (AMH) of less than or equal to 5.4 pmol/l indicates that a poor response is likely, whereas more than or equal to 25 pmol/l is likely to give a good response.
- If serum FSH (early in the cycle) is more than 8.9 iu/l, a poor response is anticipated, compared to a good response if less than 4 iu/l.

WHO classifies ovulatory disorders into three groups:

- **Group I** Hypothalamic pituitary failure – this may be BMI dependent
- **Group II** Hypothalamic pituitary ovarian dysfunction – which is mainly PCOS
- **Group III** Ovarian failure – end organ failure

**Tubal factors**

The question is whether the tubes are patent to allow the sperm to travel to meet the ovum. This can be answered either by an X-ray in the form of a hysterosalpingogram (HSG) (Fig. I.2), or by dye laparoscopy as both techniques should show whether the tubes are patent. The hysterosalpingogram does not require general anaesthesia and is recommended if there are no co-morbidities. It should be performed.
after menstruation has finished and will outline the cavity of the uterus. More recently, hysterosalpingo contrast sonography (HyCoSy) has been recommended if the expertise is available. The dye laparoscopy will show the tubes and any spill of dye. It needs to be performed under a general anaesthetic and is recommended where co-morbidities are anticipated (e.g. endometriosis). It will not tell anything about the uterine cavity.

Other investigations
There are a number of investigations that are no longer considered to be of value and are listed here for completeness only:

- Antisperm antibodies
- Hysteroscopy and endometrial biopsy should only be performed if indicated
- Post-coital test is no longer recommended and of no predictive value

Male factors
The causes of male infertility fall into the following broad categories, which may lead to oligospermia, necrospermia or azoospermia:

- Sperm production problems. These include:
  - Chromosomal or genetic causes. Men suffering from the chromosomal anomaly Klinefelter’s syndrome (XXY) have undeveloped genitalia with small, soft testes, and they are infertile.
  - Undescended testes (failure of the testes to descend at birth). Cryptorchidism may affect the function of the testis and make it more likely that a malignancy will develop within it. The testis in this case is almost certainly sterile.
  - Infections including gonorrhoea can block the epididymis, and mumps can cause atrophy of the testes if the individual suffers orchitis.
  - Heat.
  - Varicocoele, which may increase the local temperature of the testis. The presence of a varicocoele may affect sperm production and removal may be warranted.
  - Drugs and chemicals.
  - Radiation damage.
  - Unknown.
- Sperm DNA damage or fragmentation. This includes:
  - Male age over 40 years.
  - Long periods of abstinence.
  - Lifestyle factors (e.g. smoking nicotine and especially marijuana, alcohol, poor diet and medications, which include sulfasalazine, chemotherapeutic agents, antimicrobials and some antihypertensives).
  - Sexual problems, including:
    - Retrograde and premature ejaculation.
    - Failure to ejaculate.
    - Infrequent intercourse and low libido.
    - Spinal cord injury or damage.
    - Prostate surgery.
    - Erectile dysfunction.
- Hormonal problems:
  - Pituitary tumours (hyperprolactinaemia)
  - Congenital lack of LH/follicle-stimulating hormone (FSH).
  - Anabolic steroid abuse.

Female factors
Fertility problems occur in a number of areas:

- Sexual dysfunction: includes dyspareunia or vaginismus leading to infrequent coitus.
- Cervical factors: cervical lesions include abnormalities of cervical secretion and stenosis of the cervix following treatment for premalignant disease. These are uncommon.
- Uterine and/or tubal problems: gross pelvic lesions, which include absence of the uterus, vagina, Fallopian tubes or ovaries, closure of the hymen or vagina, fibroids, polyps, carcinoma, tuberculosis of the endometrium and endometriosis. Tubal lesions include inflammatory disease due to infections e.g. Chlamydia and tuberculosis. Occasionally, the tubes may be rudimentary.
- Endocrine problems, these are classified earlier as ovulatory disorders (Groups I–III) but may include problems with other endocrine glands which would need rectifying.
- Previous obstetric history. Injury/infection/postpartum haemorrhage leading to pituitary problems.
- Contraception use: oral contraceptive pill or intrauterine contraceptive device.
- Constitutional: chromosomal anomalies such as Turner’s syndrome (XO) and super-female status (XXX), anxiety and also, for completeness, anaemia and age.
- Age: fertility is problematic with increasing age of the woman, as previously discussed.
JAUNDICE

Simon Anderson

TYPES
Jaundice may be caused by a raised conjugated or unconjugated bilirubin. Unconjugated hyperbilirubinaemia may be due to excessive production of bilirubin (haemolysis), reduced uptake of bilirubin, or a failure of conjugation by the liver. Conjugated hyperbilirubinaemia results from hepatocellular damage or obstruction of the bile ducts, either within the liver (intrahepatic cholestasis) or of a major bile duct (extrahepatic obstructive jaundice). Jaundice is also often classified into pre-hepatic (haemolytic), hepatic and extrahepatic (obstructive) types. Clues to the cause of jaundice may be obtained from the history and physical examination (Tables J.1 and J.2).

INVESTIGATIONS OF JAUNDICE
The simplest investigations are liver function tests and urine examinations; typical abnormalities are shown in Table J.3. The liver enzymes aspartate transaminase (AST) and alanine transaminase (ALT) are normally contained within the liver cells and are released during hepatocellular necrosis, whereas alkaline phosphatase (ALK-P) and glutamyl transferase (GGT) are excreted into the biliary system and rise in situations of intrahepatic or extrahepatic biliary stasis.

There is no bilirubin present in the urine of patients with pre-hepatic (haemolytic) jaundice, because unconjugated bilirubin is tightly bound to albumin and is not filtered at the glomerulus. On the other hand, conjugated bilirubin is water-soluble, and it stains the urine dark in hepatocellular and obstructive jaundice. Urobilinogen is produced by bacteria in the gut and is normally partially reabsorbed into the portal vein, taken up by hepatocytes and re-excreted in bile. When the liver is damaged, hepatic extraction is less efficient, and the concentration of urobilinogen in plasma, and hence in the urine, rises. The presence of urobilinogen in the urine is thus a test of liver function, and one of the earliest signs of recovery from hepatocellular jaundice is the disappearance of urobilinogen from the urine as it is again removed by the liver. In complete obstructive jaundice, urobilinogen is absent from the urine as there is no bilirubin in the gut.

All patients with jaundice should undergo an ultrasound examination of the liver which is accurate at determining whether or not there is biliary obstruction. If equivocal, the ultrasound should be repeated as jaundice deepens, and it may then be obvious that there is indeed an obstructive cause. The radiologist

<table>
<thead>
<tr>
<th>Table J.1 History of jaundice</th>
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<tr>
<td><strong>Haemolytic</strong></td>
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<tr>
<td>Family history</td>
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<tr>
<td>Racial origin</td>
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<tr>
<td>Drug history</td>
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<tr>
<td>Symptoms of anaemia</td>
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<table>
<thead>
<tr>
<th>Table J.2 Physical signs associated with jaundice</th>
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<tr>
<td><strong>Haemolytic</strong></td>
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<tr>
<td>Splenomegaly, reduced stature</td>
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May also show liver lesions, dilatation of the gallbladder or gallstones (cholelithiasis). Furthermore, the pancreas, portal and hepatic veins and the spleen can be visualized. If extrahepatic biliary obstruction is seen, endoscopic retrograde cholangiopancreatography (ERCP) should be performed. This entails the passage of a side-viewing endoscope into the second part of the duodenum (Fig. J.1). Ampullary tumours can be identified and biopsied. The bile and pancreatic ducts can be cannulated and opacified using radiological contrast material. The procedure is successful in approximately 90 per cent of patients and, as well as defining the cause of the obstruction, common bile duct stones can be removed and stents inserted to relieve obstruction. If an ERCP is unsuccessful, a percutaneous transhepatic cholangiogram (PTC) can be performed. The bile ducts are accessed by percutaneous cannulation through the liver using ultrasound or CT guidance.

A rational approach to the investigation of jaundice is illustrated in Figure J.2. Liver biopsy confirms the presence of hepatocellular damage, but it is not usually employed because of the the risks it involves (pain, bleeding, bile leak) and the accuracy of other, non-invasive investigations.

**UNCONJUGATED HYPERBILIRUBINAEMIA**

This condition is due to:

1. Increased production of bilirubin
   - Inefficient marrow production
   - Increased breakdown (haemolysis)
     - Haemoglobinopathies (e.g. sickle-cell disease)
     - Antibody-mediated
     - Drug-induced
2. Decreased uptake of bilirubin into the liver: Gilbert’s disease
3. Decreased conjugation of bilirubin in the liver
   - Crigler–Najjar syndrome
   - Neonatal jaundice
   - Drugs
   - Lucey–Driscoll syndrome

**INCREASED PRODUCTION OF BILIRUBIN WITHOUT HAEMOLYSIS (SHUNT HYPERBILIRUBINAEMIA)**

Very rarely, inefficient marrow production of haemoglobin results in increased amounts of unconjugated bilirubin being released into the circulation (‘early label’ bilirubin). The red cells manufactured, however, have a normal lifespan.
This is a rare primary condition, and it also occurs in a number of other causes of inefficient erythropoiesis, for example anaemia.

**INCREASED PRODUCTION OF BILIRUBIN DUE TO HAEMOLYSIS**

Most commonly, unconjugated hyperbilirubinaemia results from haemolysis, either caused by an intrinsic abnormality of the red cells or due to the development of an abnormal mechanism of destruction (extrinsic). The cause is usually found with basic investigations, but detailed investigations are sometimes needed and are listed below.

- **Evidence of intravascular haemolysis**
  - Haptoglobins
  - Haemoglobinaemia
  - Haemoglobinuria
  - Haemosiderinuria
  - Methaemalbuminaemia
- **Evidence of increased marrow production**
  - Reticulocytosis
  - Skeletal changes
  - Marrow hyperplasia
- **Evidence of red cell damage**
  - Fragmented forms
- **Evidence of shortened red cell lifespan**
  - Radioactive labelling of red cells

**Evidence of intravascular haemolysis**

Haemoglobin released during intravascular haemolysis is normally attached to haptoglobin, the levels of which are usually reduced in chronic haemolytic states. However, levels may also be reduced by chronic liver disease and increased non-specifically in a number of connective tissue disorders. Haemoglobinaemia (sometimes associated with methaemalbuminaemia), haemoglobinuria and haemosiderinuria provide incontrovertible evidence of intravascular haemolysis, but are frequently absent in chronic haemolytic anaemias.

**Evidence of increased marrow production**

In a compensated anaemia, reticulocytosis with a raised mean corpuscular volume is common. Increased marrow activity results in skeletal changes, which are frequent in thalassaemia and sickle-cell disease but rare in other conditions. The skull of such patients demonstrates a thickened vault, and the diploe are widened. Bony trabeculae arising at right angles to the diploe may produce a ‘hair-on-end’ appearance. The bones of the limbs have a widened narrow cavity with a coarse trabecular pattern.

**Evidence of red cell damage**

Fragmented forms may provide evidence of increased red cell destruction.

**Evidence of shortened red cell lifespan**

The standard clinical test to detect shortened red cell survival is to tag the patient’s cells with radioactive chromium and to measure the decline in plasma radioactivity.

**INTRINSIC DEFECTS OF THE RED CELLS LEADING TO INCREASED HAEMOLYSIS**

These include:

- Spherocytosis
- Elliptocytosis
- Enzyme defects
- Haemoglobinopathies
  - Sickle-cell disease
  - HbSC disease
  - Thalassaemia
- Paroxysmal nocturnal haemoglobinuria

**Congenital spherocytosis**

This is a dominantly inherited defect that probably affects the red cell membrane, rendering it more permeable to sodium. The red cells are spherical rather than the usual biconcave shape, and they are more readily haemolysed in hypotonic saline (red cell fragility test). The patient usually presents in early childhood, even though jaundice may have been noticed in early childhood; often, jaundice is first identified in the teens. Splenomegaly is common; bile pigment stones are frequently formed, and patients occasionally present with obstructive jaundice. The disease is characterized by crises of worsening anaemia and jaundice caused by increased haemolysis due to infection. The diagnosis is usually straightforward, with splenomegaly, a family history and a typical blood film, but it must be remembered that spherocytes may be a feature of a number of different types of haemolytic anaemia, and that occasional mild cases of congenital spherocytosis do not present until adulthood. Splenectomy is usually followed by long-term remission of symptoms.

**Congenital elliptocytosis**

This is another Mendelian dominant disorder, and it is usually asymptomatic, without haemolysis. Occasionally, a compensated haemolytic anaemia occurs, but anaemia sufficient to produce jaundice is extremely rare.

**Enzyme defects**

A wide variety of enzyme defects in the red cells have been described which produce a haemolytic anaemia and jaundice. These are usually recessively inherited. Suspicion of this type of disorder is always aroused if haemoglobin electrophoresis and osmotic fragility...
in vitro sickling haemoglobin, and by demonstrating the which will show increased amounts of fetal and A2

The diagnosis is made by haemoglobin electrophoresis, still dying either as children or in young adult life. jaundice. The prognosis is serious, with many patients are again common, which may produce obstructive is common in children, repeated infarcts of the spleen and bone pain and a high fever. Although splenomegaly in the circulation, which produces severe abdominal frequently punctuated by crises of spontaneous sickling clinical manifestations are variable, life for the patient is those of African origin, and it is characterized by jaundice, anaemia and skeletal changes. Although the clinical manifestations are variable, life for the patient is usually only mildly affected and do not become jaundiced. Patients who inherit two sickle-cell genes have sickle-cell disease. An amino acid substitution in the beta chain produces an unstable haemoglobin molecule, which polymerizes into the reduced state, producing long chains that distort the red cells into a sickle shape. Sickle-cell disease predominantly affects those of African origin, and it is characterized by jaundice, anaemia and skeletal changes. Although the abnormal haemolysis of the red cells can be demonstrated if the plasma is acidified (Ham's test). Paroxysmal nocturnal haemoglobinuria Episodes of haemolysis may be accompanied by slight jaundice in this rare condition. Diagnosis can be made by the characteristic history of red urine, which contains haemoglobin, following sleeping. The abnormal haemolysis of the red cells can be demonstrated if the plasma is acidified (Ham's test).

EXTRINSIC FACTORS LEADING TO INCREASED HAEMOLYSIS These include:

- Autoimmune haemolytic anaemia
- Cold haemoglobinuria
- Drugs and chemicals
- Glucose-6-phosphate dehydrogenase deficiency
- Miscellaneous

Warm antibody autoimmune haemolytic anaemia In this condition, antibody coats the patient’s red cells at 37 °C, and this results in increased
extravascular destruction. The antibody is usually of the IgG class and incomplete (i.e. does not directly cause agglutination or haemolysis). This antibody can be detected by the direct Coombs’ test, where the addition of antibody to IgG in vitro causes the red cells to agglutinate. Spherocytes are often present in the blood film, the white count may be raised, and occasionally the platelets are low, producing purpura. In acute acquired haemolytic anaemia, the patient is usually a child with a palpable spleen, jaundice, anaemia and the constitutional symptoms of fever, vomiting and prostration. In the chronic form, the onset is insidious, usually in adults, but again the patient is usually jaundiced (in 75 per cent of cases) and has a palpable spleen. In half the patients with acquired haemolytic anaemia, no cause for the antibody formation is found, but in the remainder it is secondary to a number of diseases, most importantly systemic lupus erythematosus, but also recticuloendothelial malignancy, leukaemia and sarcoidosis. Resolution of the symptoms usually occurs following treatment with corticosteroids or, occasionally, splenectomy.

**Cold antibody autoimmune anaemia**

Occasionally, antibody is produced that reacts with the patient’s red cells at low temperatures. Depending on the thermal range of the antibody, a continuous mild haemolytic anaemia may occur that is punctuated by cold agglutinins. It may also follow a mismatched blood transfusion or occur in a severely burned patient. Excessive exercise, particularly on hard roads, may also lead to episodes of intravascular haemolysis (march haemoglobinuria).

**Drug and chemical-induced haemolysis**

Some chemicals (e.g. arsenic and naphthalene in mothballs) produce haemolysis and jaundice that is directly dose-related. Haemolysis may occur with other drugs; this is unrelated to the dose, and occurs only in a few susceptible individuals (Box J.1). The two important mechanisms for producing haemolysis in this situation are an associated deficiency of glucose-6-phosphate dehydrogenase in the red cell, or the production of autoantibodies, often directed against the drug attached to the red cell membrane, which acts as a hapten. In this type of haemolysis, the blood film often shows spherocytes and red cell inclusions (Heinz bodies).

**Glucose-6-phosphate dehydrogenase deficiency** is common in black individuals and inhabitants of the Mediterranean littoral. This enzyme helps to maintain the cell concentration of reduced glutathione, which in turn stabilizes the haemoglobin molecule.

**Favism** is a disorder characterized by intravascular haemolysis and jaundice occurring when a glucose-6-phosphate dehydrogenase-deficient patient – usually a child from the Mediterranean region – ingests fava beans or inhales the pollen.

Similar episodes occur when the patient is exposed to certain drugs (see Box J.1). Haemolysis ceases when the older population of red cells containing less of the enzyme is destroyed.

**Autoimmune drug-induced haemolysis** is particularly common with methyldopa, where the direct Coombs’ test is positive in 20 per cent of patients taking the drug, but haemolysis occurs in less than 1 per cent.

**Other conditions causing haemolysis**

Acute haemolysis with jaundice may occur with various infections (e.g. malaria and gangrene), and more mildly with viral pneumonia (usually due to cold agglutinins). It may also follow a mismatched blood transfusion or occur in a severely burned patient.

Excessive exercise, particularly on hard roads, may also lead to episodes of intravascular haemolysis (march haemoglobinuria).

**Microangiopathic haemolytic anaemia** is the name given to a group of conditions characterized by haemolysis in association with fragmentation of red cells as they pass through blood vessels damaged by clots. Evidence of disseminated intravascular coagulation is common. Such haemolytic anaemias are present in thrombotic thrombocytopenic purpura, malignant hypertension, disseminated neoplasia and in association with uraemia in children (the haemolytic-uraemic syndrome).

**UNCONJUGATED HYPERBILIRUBINAEMIA CAUSED BY IMPAIRED UPTAKE OF BILIRUBIN INTO THE LIVER**

**Gilbert’s disease**

There is some debate as to whether this condition exists, or whether it represents the upper range of
a normal population distribution of unconjugated bilirubin. However, most would accept that it is a common familial condition in which there is a mild degree of unconjugated hyperbilirubinaemia. It is probably inherited as a Mendelian dominant with variable penetrance. The degree of jaundice varies and often increases following an infection or a period of fasting. Episodes of deepening jaundice may be associated with recurrent, vague abdominal pains. Mild decreases in red cell lifespan and the liver’s ability to conjugate bilirubin are associated with a failure of transport of unconjugated bilirubin into the liver cell. The diagnosis is made by excluding liver disease. It may be confirmed by fasting the patient or by giving an intravenous injection of nicotinic acid. Both of these manoeuvres result in an increase in serum bilirubin concentrations. They are rarely necessary, however, since the diagnosis is usually obvious.

**UNCONJUGATED HYPERBILIRUBINAEMIA CAUSED BY IMPAIRED CONJUGATION IN THE LIVER**

**Crigler–Najjar syndrome**
In this familial condition, there is a deficiency of the liver enzyme glucuronyl transferase, which conjugates bilirubin. In severely affected patients (type 1), death occurs during the neonatal period. A partial enzyme defect with some conjugated bilirubin in the bile and a better prognosis (type 2) also occurs.

**Neonatal jaundice**
Glucuronyl transferase matures shortly before birth, and newborn babies – particularly if premature – will become mildly jaundiced. This may be severe in conditions increasing the bilirubin load (e.g. haemolysis), and kernicterus may result. Occasionally, prolonged neonatal jaundice is thought to occur in breastfed babies due to the presence of pregnanediol in the milk.

**Lucey–Driscoll syndrome**
Unconjugated bilirubin has rarely been described in pregnancy due to hormonal inhibition of bilirubin conjugation.

**CONJUGATED HYPERBILIRUBINAEMIA**
The hepatic causes of conjugated bilirubinaemia may be divided into: (i) acute hepatocellular damage, associated with considerable increases in hepatic enzymes and a short clinical course; and (ii) chronic damage, where the course is protracted and there are lesser rises in liver enzymes:
- Acute liver damage
  - Viral hepatitis
- Non-viral hepatitis
- Drug-induced
- Poisons
- Fatty liver of pregnancy
- Chronic liver damage
  - Cirrhosis
  - Tumours
    - Primary
    - Secondary
- Infections
  - Reticuloendothelial tumours
  - Amyloidosis

**Acute hepatic damage**
Causes of acute hepatic damage may produce a mild clinical illness or a severe disease (fulminant hepatic failure) with encephalopathy and a high mortality, when cerebral oedema, renal failure and a bleeding diathesis are frequent causes of death.

**Viral hepatitis**
Although a large number of viruses occasionally cause hepatitis (including rubella, Coxsackie B, herpes simplex, yellow fever virus and cytomegalovirus), the four common ones are virus A, virus B, virus C and infectious mononucleosis. The recognition of serological markers for hepatitis A and B has revolutionized our understanding of viral hepatitis, and it is now realized that many patients with these infections do not become jaundiced.

**Virus A (infectious hepatitis)**
This is an endemic infection that is caused by a 27 nm RNA virus with a short incubation period (15–20 days). Outbreaks usually occur in conditions of poor hygiene or overcrowding. The usual transmission is faeco-oral. Patients present with malaise, anorexia, fever and a rapid onset of jaundice. There is often a vague ache in the right upper quadrant, and the liver is enlarged and tender. Complete recovery is usually within a few weeks, although relapses may occur. Occasionally, patients develop deep jaundice due to intrahepatic cholestasis during the convalescent period. The diagnosis is confirmed by the demonstration of IgM antibody to the virus.

**Virus B (serum hepatitis)**
This infection has a longer incubation period, and arthralgias and rashes may occur in the prodromal period. It is caused by a DNA virus that has an outer coat derived from the host cells and an inner core. Originally, only blood transmission was recognized (e.g. blood transfusions, transfusions of blood products; haemophilic globulin), or by needles contaminated with blood in drug addicts and in tattooing. Outbreaks in renal dialysis units produced by blood contamination have caused great concern.
in the past. The disease is also venereally transmitted, as the virus is present in semen. Hepatitis B is common in homosexuals, 10 per cent of whom have serological markers of past infection. Asymptomatic carriers of hepatitis B infection are very frequent in certain parts of the world (e.g. Africa, China and parts of the Mediterranean). These patients may transmit the infection vertically from mother to offspring. The diagnosis of virus B hepatitis is made by the detection of the presence of surface antigen (HbsAg) in the bloodstream. For the patient to be infectious, whole virus (Dane) particles must be present in the bloodstream, and the presence of e antigen is a marker for this.

The majority of patients develop an acute viral hepatitis, and this is associated with formation of antibodies and clearance of the virus from the liver. Other individuals fail to clear the virus and become carriers, some of these developing chronic liver disease. Patients with persistent hepatitis B virus infection are at risk of developing primary hepatocellular carcinoma.

Hepatitis D virus is an RNA virus that replicates in patients with acute or chronic hepatitis B infection. Simultaneous infection with both B and D virus leads to an acute exacerbation of hepatitis, which may be self-limiting. When chronic infection develops with co-infection, the hepatitis is frequently more severe.

Hepatitis C The prevalence of hepatitis C is approximately 2–3 per cent worldwide, with a higher prevalence in certain areas such as the Middle East, where rates may be as high as 14 per cent. Hepatitis is associated with an increased risk of cirrhosis and hepatocellular carcinoma. The mode of transmission is similar to that of hepatitis B. The hepatitis C virus is a small, enveloped RNA virus. In acute hepatitis C, the clinical course may be mild, only 25 per cent of patients become jaundiced, and these patients are more likely to clear the virus. Most cases of chronic hepatitis C are not preceded by an acute illness. Serum transaminases remain elevated in 60–85 per cent of patients at 1 year after diagnosis, and there is histological evidence of chronic hepatitis. Cirrhosis develops in approximately 10 years in 10–20 per cent of cases with chronic disease, and hepatitis C RNA persists in patients with abnormal liver enzymes.

Hepatitis E is an RNA virus, of which there are many genotypes. It is endemic in Asia, Africa, the Middle East and Central America, and is associated with poor sanitation. The seroprevalence is approximately 5 per cent in children aged under 10 years, and 10–40 per cent in adults. The virus is particularly virulent in pregnant women, among whom there is a high fatality rate of approximately 20 per cent. In non-endemic regions, hepatitis E accounts for less than 1 per cent of cases of acute viral hepatitis.

**Infectious mononucleosis** Up to 15 per cent of patients with glandular fever develop jaundice. The clinical picture is characteristic, with malaise, sore throat, skin rashes, lymphadenopathy and splenomegaly. Atypical mononuclear cells are found in the peripheral blood, and the test for heterophile antibody (Paul-Bunnell) is usually positive.

**Yellow fever** This is a zoonosis and is transmitted to man from a primate pool by the mosquito in tropical Africa, the Caribbean and South America. The incubation period is short (3–4 days), with a sudden onset of rigors, jaundice and abdominal pain. Its course may be fulminant with renal failure and a bleeding diathesis.

**Non-viral infections**

**Relapsing fever** This condition is caused by a spirochaete of the *Borrelia* group of bacteria and is characterized by jaundice and a fever of up to 40°C, which normally lasts for 4–5 days and then remits. The epidemic form of the disease is usually caused by lice and is common during periods of famine. An endemic infection is usually transmitted by the tick, and it is common in the Far East, Africa and America.

**Leptospirosis** The spirochaete *Leptospira icterohaemorrhagica* infects a variety of small animals, and humans contract the disease by bathing in water contaminated with these infected animals’ urine. The disease is biphasic, with an initial illness a few days after exposure, a temperature, meningism and prominent myalgias and conjunctivitis. Recovery may occur, or after a week the patient may develop widespread bruising, jaundice and occasionally renal failure. *Leptospira icterohaemorrhagiae* is transmitted by rats’ urine, mainly to agricultural and sewage workers, and produces a severe form of the disease where jaundice and renal failure are particularly likely to occur.

**Other bacterial infections** Jaundice may complicate any septicaemic illness. Occasionally, an infected thrombus in the portal vein may occur (portal pyaemia) following an acute infection in the area drained by the portal system (e.g. appendicitis). The signs of a portal pyaemia are severe prostration, a swinging pyrexia and jaundice.

**Drug-induced acute hepatic damage**

Drugs either produce predictable dose-related hepatic necrosis (e.g. paracetamol) or, more commonly, damage...
is produced unpredictably in only a few of the patients exposed to the drug and unrelated to its dosage. There are two basic patterns of liver damage: acute hepatic cellular necrosis with features identical to viral hepatitis, and intrahepatic cholestasis. It is not possible to provide an exhaustive list of drugs producing hepatocellular damage (Table J.4), and a high index of suspicion should exist in any jaundiced patient who is taking drugs. There are certain situations that will make the individual more susceptible to liver damage; for example, reactions are more likely at the extremes of age, HIV infection renders the liver more sensitive to co-trimoxazole and sulphonamides, enzyme inducers increase sensitivity to rifampicin and isoniazid, enzyme inhibitors to oestrogens, while in any condition that interferes with nutrition (e.g. intravenous drug or alcohol abuse, or HIV infection), malabsorption increases the toxicity of paracetamol.

The website: livertox.nih.gov provides a useful comprehensive guide to drug-induced hepatotoxicity.

### Table J.4 Drug-induced hepatic damage (drugs in italics are the most common causes of liver damage)

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<tr>
<th>Drug</th>
<th>Acute hepatic necrosis</th>
<th>Cholestasis</th>
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<tr>
<td>Paracetamol</td>
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<tr>
<td>Dextropropoxyphene</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>Halothane</td>
<td>+</td>
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</tr>
<tr>
<td>Tetracycline</td>
<td>+ (in pregnancy)</td>
<td></td>
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<tr>
<td>Erythromycin estolate</td>
<td>+</td>
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<tr>
<td>Penicillin</td>
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<tr>
<td>Sulphasalazine</td>
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<tr>
<td>Nitrofurantoin</td>
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<tr>
<td>Pheniramine maleate</td>
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<tr>
<td>Piperazine</td>
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<tr>
<td>Piperazine</td>
<td>+</td>
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<tr>
<td>Rifampicin</td>
<td>+</td>
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<tr>
<td>Para-aminosalicylic acid</td>
<td>+</td>
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<tr>
<td>Chlorpromazine</td>
<td>+</td>
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<tr>
<td>Monoamine oxidase inhibitor</td>
<td>+ (cirrhosis)</td>
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<tr>
<td>Methyldopa</td>
<td>+</td>
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<tr>
<td>Quinidine</td>
<td>+ (cirrhosis)</td>
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<tr>
<td>Perhexiline</td>
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<tr>
<td>Chlorpropamide</td>
<td>+</td>
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<tr>
<td>Phenytoin</td>
<td>+</td>
<td>+</td>
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<tr>
<td>Propylthiouracil</td>
<td>+</td>
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<tr>
<td>Oral contraceptives</td>
<td>+</td>
<td></td>
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<tr>
<td>Anabolic steroids</td>
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</tbody>
</table>

**Paracetamol** Paracetamol overdose is the most common cause of fulminant hepatic failure in the UK. Hepatic damage is dose-related, but death has been reported with amounts as low as 7.5 g. Following ingestion, paracetamol is metabolized to a toxic intermediate, which is scavenged by glutathione. When glutathione stores are exhausted, the metabolite binds covalently to the membrane of hepatocytes, causing cell death. Chronic alcoholics whose microsomal enzymes are reduced and whose glutathione stores tend to be depressed are at increased risk following the overdose. Nausea, vomiting and abdominal pain develop within 12–36 hours, and jaundice develops 2–3 days later. In severe cases, this leads to liver failure with coagulopathy and encephalopathy. A very characteristic feature of paracetamol poisoning is the development of renal failure.

**Halothane** Halothane is a very safe anaesthetic, but there is a small incidence (0.003 per cent) of serious acute hepatic necrosis following the use of the drug, which may lead to fever, jaundice and death. Inadvertent repeated use in patients who were previously jaundiced following exposure to halothane results in recurrence of the patient’s jaundice. Halothane hepatitis is more common after repeated exposures, particularly in obese patients. Jaundice associated with pyrexia usually occurs 2 weeks after the initial exposure, but only 10 days after subsequent administration. Halothane should not be re-used in patients who have suffered a febrile illness and abnormal liver function tests after a previous encounter with this agent.

**Oral contraceptives** The older oral contraceptives containing a relatively high concentration of oestrogen occasionally led to a mild cholestatic jaundice. These individuals are particularly prone to develop cholestasis in pregnancy. In addition, the older contraceptives also had a tendency to cause Budd–Chiari syndrome and a variety of tumours within the liver, in particular benign adenomas. The newer contraceptives, which contain much lower concentrations of oestrogens, are much safer and rarely cause these complications.

**Anabolic steroids** The C17-alpha-alkylated-substituted testosterones (e.g. norethisterone and norethanandrolone) produce a dose-related cholestasis by a similar mechanism to the oral contraceptive.

**Chlorpromazine** An unpredictable cholestatic jaundice may occur in 1 per cent of patients within a month of starting treatment with this drug. Eosinophilia and mitochondrial antibodies are frequently found in the bloodstream. The patient itches and has pale stools.
and dark urine. Three-quarters of patients recover on withdrawal of the drug, but a few develop prolonged cholestasis resembling primary biliary cirrhosis (see below).

**Antineoplastic drugs** A variety of such drugs cause jaundice and liver damage (e.g. methotrexate). However, the primary conditions for which these drugs are administered are often a cause for jaundice.

**Industrial toxins**

These are only rarely a cause of jaundice in humans. Most seem to act by inhibiting protein synthesis. Carbon tetrachloride and (less commonly) other volatile hydrocarbons produce acute hepatocellular necrosis and jaundice within 1–2 days of exposure. Renal failure, pancreatitis, pulmonary oedema and death may also occur. Dicophane (DDT) and trinitrotoluene (TNT) also occasionally produce hepatic necrosis.

A cholestatic jaundice has been described following the accidental ingestion of flour contaminated by diaminodiphenyl methane (so-called ‘Epping jaundice’, named after the place where the outbreak occurred).

**Amanita** Ingestion of as little as three wild mushrooms of the *Amanita* species may be fatal. Abdominal pain and diarrhoea occur within 18 hours of ingestion, followed 3 days later by the development of fulminant hepatic failure and jaundice.

**Acute fatty liver of pregnancy**

This condition occurs in 1 in 15 000 cases. It is common in primigravida and those who have had multiple pregnancies. It is associated with genetically determined defects of beta-oxidation of fatty acids in some cases. Onset occurs in the third trimester, with vomiting, upper abdominal pain, anorexia and malaise. Later, disseminated intravascular coagulation, leucocytosis, hypoglycaemia and hyperammonaemia develop, and progress to fulminant hepatic failure. Mortality rates (mother and baby) are high if the condition is not recognized and is allowed to progress. Treatment is by prompt delivery of the baby.

**CHRONIC LIVER DAMAGE: CIRRHOSIS**

Cirrhosis is defined as diffuse fibrosis with nodular regeneration of the liver, which destroys the normal spatial relationship of the lobules. It is classified as micronodular, macronodular or mixed, depending on the size of the nodules. The condition may be suspected in cases of jaundice where the liver is firm and palpable, although sometimes it is shrunken and impalpable. The stigmata of chronic liver disease (see Table J.2) are often present, and the spleen may be enlarged because of portal hyper-tension. Oedema and ascites are caused by a combination of portal hypertension, sodium retention and hypoaalbuminaemia. Hepatic encephalopathy may occur, with a sweet, musty odour to the breath (caused by mercaptans originating from the gut breakdown of methionine), a flapping tremor (asterixis), disorders of the sleep rhythm, drowsiness and, in severe cases, coma. A chronic form of encephalopathy with slowness and psychiatric changes may also occur. Jaundice is a relatively late complication of cirrhosis, and many patients with compensated cirrhosis have relatively normal liver function tests.

**Causes of cirrhosis** include the following:

- Alcoholic liver disease
- Infections
  - Viral (hepatitis B and C viruses)
  - Bacterial (syphilis)
  - Protozoan (schistosomiasis)
- Chronic active hepatitis
  - Lupoid
  - Hepatitis B virus
  - Drugs
- Genetic defects
  - Haemochromatosis
  - Wilson’s disease
  - Galactosaemia
  - Glycogen storage diseases
  - Alpha-1-antitrypsin deficiency
- Biliary disease
  - Long-standing extrahepatic biliary obstruction
  - Primary biliary cirrhosis
  - Sclerosing cholangitis
  - Congenital hepatic fibrosis
- Venous congestion
  - Cardiac failure
  - Constrictive pericarditis
  - Budd–Chiari syndrome
- Jejuno-ileal bypass

**Alcoholic liver disease**

There is a strong relationship between the national figures for consumption of alcohol and death from cirrhosis. Women are more susceptible to the effects of alcohol than men, and drinking more than 20 g a day is associated with an increased risk of cirrhosis. In men, a consumption of over 80 g a day is associated with an increased risk, which rises to a 25-fold increase if more than 100 g a day is consumed. Individual susceptibility is also important – some people never develop cirrhosis, however much they drink. There are three types of liver disease associated with alcohol:

- **Fatty infiltration** does not cause jaundice and does not lead to cirrhosis.
smooth muscle antibodies, antinuclear factor and that is characterized by the presence of circulating is an autoimmune disease predominantly of women. Autoimmune chronic active hepatitis (lupoid hepatitis) presenting with hepatitis B in the Western hemisphere. to blood products is often elucidated in patients intravenous drug use, homosexuality or exposure is chronic hepatitis B virus infection. A history of diseases. The most common cause worldwide Chronic active hepatitis is a consequence of a variety e.g. hepatitis B infection, which is also very common uncommon and possibly due to associated conditions with only minor liver parenchymal injury. Cirrhosis is produced by multiple protein inclusions (Mallory’s alcoholic haemolytic anaemia and hyperlipidaemia in addition to jaundice. • Cirrhosis: alcohol produces a micronodular cirrhosis, and the prognosis is undoubtedly worse if the patient continues to drink. It has been suggested that autoimmune mechanisms, particularly in alcoholic hepatitis, may be important. A combination of alcohol abuse and hepatitis B or C infection is particularly likely to lead to cirrhosis and strongly predisposes to hepatic carcinoma.

Infections
The discovery of markers for hepatitis B and C has demonstrated that these are both important causes of cirrhosis. Congenital syphilis may produce a pericellular fibrosis, but true cirrhosis is uncommon, as regeneration nodules do not usually develop.

Schistosomiasis classically produces periportal fibrosis (pipestem fibrosis) leading to portal hypertension with only minor liver parenchymal injury. Cirrhosis is uncommon and possibly due to associated conditions (e.g. hepatitis B infection, which is also very common in these patients).

Chronic active hepatitis
Chronic active hepatitis is a consequence of a variety of diseases. The most common cause worldwide is chronic hepatitis B virus infection. A history of intravenous drug use, homosexuality or exposure to blood products is often elucidated in patients presenting with hepatitis B in the Western hemisphere. Autoimmune chronic active hepatitis (lupoid hepatitis) is an autoimmune disease predominantly of women that is characterized by the presence of circulating smooth muscle antibodies, antinuclear factor and high titres of immunoglobulins (IgG). Chronic active hepatitis may also be due to the ingestion of drugs, including, methyldopa, antituberculous agents and anticonvulsants.

Chronic active hepatitis may present with malaise, jaundice and eventually splenomegaly and hepatic decompensation. The diagnosis is suggested by a hypertransaminasemia for at least 6 months. There may be evidence of chronic hepatitis B infection, circulating smooth muscle antibodies or antinuclear factor. Liver biopsy reveals an inflammatory infiltrate radiating from the portal tracts and broaching the limiting plates. Fibrosis is almost invariable, and may encircle groups of hepatocytes, resulting in the formation of rosettes. Frank cirrhosis may be present at the time of diagnosis.

Genetic defects
Haemochromatosis
This is an autosomal recessive disease characterized by increased intestinal iron absorption. The disease is rarely presents in premenopausal women. Slate-grey pigmentation develops because of melanin and iron deposition in the skin. Cirrhosis occurs; the liver is invariably enlarged, and there is evidence of portal hypertension and hepatic decompensation. Iron deposition in the pancreas causes diabetes mellitus (‘bronze diabetes’) and in the heart a cardiomyopathy. Accumulation in other endocrine glands leads to testicular atrophy, gynaecomastia and loss of body hair; hypopituitarism can also occur. The diagnosis is suggested by a high serum ferritin concentration and confirmed by liver biopsy. Hepatocellular carcinoma is a relatively common complication and cause of death.

Wilson’s disease
Wilson’s disease is a recessively inherited disease of impaired copper metabolism. Copper accumulates in the liver, causing cirrhosis, and also in the basal ganglia of the brain, causing an extrapyramidal neurological syndrome. Patients usually present between the ages of 12 and 23 years, with neurological and psychiatric manifestations. Patients usually have few liver signs until cirrhosis and portal hypertension develop. The presentation can be acute, with fulminant hepatic failure and severe haemolytic anaemia.

The diagnosis is made by slit-lamp examination of the eyes, when Kayser–Fleischer rings can be demonstrated. Urinary 24-hour copper excretion is increased, and this increases further with penicillamine therapy. In addition, serum copper concentrations are increased, and caeruloplasmin levels are low. Liver biopsy shows cirrhosis. Copper stains are positive.
Other metabolic errors
All the other metabolic errors leading to cirrhosis and jaundice mentioned above are exceedingly rare. Among the glycogen storage diseases, only type IV leads to cirrhosis and jaundice.

Alpha-1-antitrypsin deficiency
This autosomal recessively inherited disease may present as a severe neonatal hepatitis or as established cirrhosis in patients below the age of 20 years. There may be associated lung disease, which characteristically presents as emphysema, pulmonary fibrosis and respiratory failure. Liver biopsy shows periodic acid–Schiff-positive-stained inclusion bodies within hepatocytes.

Biliary disease
Disease affecting the extrahepatic biliary tract, intrahepatic ductules or canaliculi can lead to cirrhosis. Extrahepatic biliary obstruction following trauma to the bile ducts can occasionally cause a secondary biliary cirrhosis; extrahepatic biliary obstruction is considered below. The most important causes of intrahepatic cholestasis are drugs and primary biliary cirrhosis.

Primary biliary cirrhosis
Primary biliary cirrhosis (PBC) predominantly affects middle-aged females and is due to destruction of bile ductules by a cell-mediated autoimmune process. The disease progresses slowly in the majority of patients, and usually presents with a cholestatic syndrome comprising itching, pale stools and dark urine. Hypercholesterolaemia is common and may lead to xanthelasmata. The disease is associated with other autoimmune diseases, including hypothyroidism, Addison’s disease, systemic sclerosis, diabetes and renal tubular acidosis. Some patients present with bleeding oesophageal varices or ascites, without a history of itching.

PBC should be considered in any women with cholestatic liver function tests. The mitochondrial antibody is positive in more than 98 per cent of patients. Serum IgM concentrations are increased. A liver biopsy shows chronic inflammatory infiltrate in the portal tract, with destruction and paucity of the bile ductules. Granulomas may be seen within the portal tracts.

Sclerosing cholangitis
Secondary sclerosing cholangitis (PSC) describes fibrosis and narrowing of the intra- and extrahepatic biliary tree due to chronic biliary sepsis, usually as a consequence of bile duct injury. Primary sclerosing cholangitis is disease of unknown aetiology of progressive destruction and fibrosis of the biliary tree. Most (80 per cent) of cases are associated with ulcerative colitis (usually mild or asymptomatic). Patients may have no liver symptoms at all or present with cholestatic liver function or fluctuating jaundice. Patients are at increased risk of both cholangiocarcinoma and colorectal cancer and need to be screened regularly for these conditions. Dilatation of dominant strictures at ERCP can provide short-term relief, but liver transplantation is the only effective treatment.

Hepatic congestion
True cirrhosis due to heart failure is very uncommon (cardiac cirrhosis), although jaundice may occur in association with chronic right heart failure.

Budd–Chiari syndrome
This occurs predominantly in women in their 20s and 30s due to occlusion to one or more of the hepatic veins (see LIVER, ENLARGEMENT OF, p. 347). Patients usually have an undetected procoagulant state (myeloproliferative disease, oral contraception) and present with hepatomegaly and ascites – jaundice being an infrequent late sign. The liver scan often shows a central area of uptake, which is due to the enlargement of the caudate lobe of the liver, the veins of which drain separately into the inferior vena cava (IVC).

CHRONIC LIVER DAMAGE: INFILTRATIONS
Amyloidosis
The liver may be involved by infiltration with amyloid, both in the primary disease, and where it is secondary to chronic suppuration, myelomatosis or rheumatoid arthritis. Amyloid is an antigen–antibody complex that characteristically shows green birefringence with Congo red staining. The liver is enlarged and rubbery, and the patient may show other features of amyloidosis, such as nephrotic syndrome, cardiac failure and malabsorption. Hepatocellular failure – and hence jaundice – is rare in this condition, and the diagnosis may be made by liver or rectal biopsy.

CHRONIC LIVER DAMAGE: TUMOURS
Metastases are 25 times more common than primary cancers and both are causes of jaundice, which carries a very poor prognosis.

Primary hepatocellular carcinoma
Hepatocellular cancer is a well recognised consequence of cirrhosis, and is found in 50–60 per cent of post-mortem examinations in cirrhotic patients. It is particularly common in certain parts of the world, especially Africa and China, where hepatitis B is the most important aetiological agent. Hepatitis C and
alcoholic cirrhosis are important causes in the West. Aflatoxin, produced by a fungus growing on grain stored in humid conditions, and the now obsolete radiocontrast material Thorotrast (thorium dioxide) are other causes.

Hepatocellular carcinoma may occur at any age, is fivefold more common in males than females, and should be suspected in any patient with cirrhosis who deteriorates or who develops a lump in the liver. A friction sound or bruit is occasionally heard over the tumour. The diagnosis is made by the finding of an elevated alpha fetoprotein and, a mass lesion in a cirrhotic liver on CT or MRI.

Primary sarcoma of the liver and malignant haemangiosarcoma

These are extremely rare tumours, which may cause jaundice in their terminal stages. Malignant haemangiosarcoma is associated with exposure to Thorotrast and vinyl chloride.

Secondary tumours

The liver is the most frequent site of blood-borne metastases, whether drained by the systemic or portal veins. It is involved in about one-third of all cancers, including half of those in the stomach, large bowel, breast and lung. The liver may be either normal in size or grossly enlarged, with palpable hard deposits. Jaundice may be absent and is usually mild. The serum alkaline phosphatase level is often markedly raised.

Reticuloendothelial diseases in the liver

The reticuloendothelial cells of the liver may be involved by a lymphoproliferative or myeloproliferative malignancy. Jaundice is usually mild, and may occasionally be due to haemolysis. Overt jaundice carries a very poor prognosis.

INTRAHEPATIC CHOLESTASIS

Causes of this condition include:

- Drugs (e.g. anabolic steroids, tricyclic antidepressants, tetracycline)
- Viral hepatitis
- Cirrhosis (occasionally)
- Dubin–Johnson syndrome
- Pregnancy
- Sclerosing cholangitis
- Biliary atresia
- Recurrent idiopathic cholestasis

In this group of conditions, the patient presents with painless obstructive jaundice, usually with pale stools, dark urine and itching, but investigations reveal no obstruction of the extrahepatic bile ducts.

Dubin–Johnson and Rotor syndromes

These are rare familial benign intermittent conditions that produce jaundice with conjugated hyperbilirubinaemia. In the Dubin–Johnson type, the liver is greenish-black and contains brown pigment. Jaundice is rarely deep, and the alkaline phosphatase remains normal. The diagnosis may be made using a bromsulphthalein (BSP) retention test, in which, after an initial fall, the serum level of BSP rises after 2 hours and remains detectable for 48 hours. The condition is thought to be due to a poor transport of conjugated bilirubin into the biliary canaliculi. The Rotor syndrome resembles Dubin–Johnson both clinically and biochemically, the main difference being the absence of brown pigment in the liver.

Pregnancy

Some women develop intrahepatic cholestasis during the last trimester of pregnancy; it is associated with itching, pale stools or dark urine. The mechanism seems similar to that of oral contraceptive-induced cholestasis.

EXTRAHEPATIC BILIARY OBSTRUCTION

Extrahepatic biliary obstruction can be classified as being due to diseases within the lumen of the bile ducts, those affecting the wall of the ducts and diseases compressing the duct from outside (Box J.2).

Obstruction is usually followed by dilatation of the common bile duct, although this may take some time to develop. The architecture of the liver is usually normal, but biopsies show pigmentation, bile plugs and infarcts. Cirrhosis can rarely develop in extremely long-standing obstruction. The urine is dark and the stools pale, clay-coloured and bulky because of their increased fat content. Biochemical investigations reveal a raised serum alkaline phosphatase, a prolonged prothrombin time because of vitamin K malabsorption, and hypocalcaemia because of vitamin D malabsorption.

### Box J.2 Causes of obstruction to the bile ducts

<table>
<thead>
<tr>
<th>Causes within the lumen of the bile ducts</th>
<th>Causes compressing the bile duct or invading it from the outside</th>
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</thead>
<tbody>
<tr>
<td>• Gallstones</td>
<td>• Pancreatic cancer</td>
</tr>
<tr>
<td>• Parasites</td>
<td>• Enlarged portal lymph nodes</td>
</tr>
<tr>
<td>• Blood (haemobilia)</td>
<td>• Aneurysm of the hepatic artery</td>
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<tr>
<td>Causes affecting the wall of the ducts</td>
<td>• Pancreatic or hydatid cysts</td>
</tr>
<tr>
<td>• Accidental division</td>
<td>• Acute or chronic pancreatitis</td>
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<tr>
<td>• Carcinoma of the bile duct (cholangiocarcinoma)</td>
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<tr>
<td>• Primary sclerosing cholangitis</td>
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</table>
The serum albumin concentration is usually maintained until the late stages.

The investigation of obstructive jaundice is described above.

**Causes due to obstruction of the bile duct lumen**

**Gallstones**

This is the most common cause of extrahepatic biliary obstruction in western countries and occurs when gallbladder stones move into the common bile duct (CBD). Occasionally, stones can form in the CBD itself in patients with a previous cholecystectomy. Patients present with severe right upper quadrant pain radiating to the back. Fever may occur if secondary (ascending) cholangitis occurs. The jaundice fluctuates and may disappear if the calculus passes into the duodenum. Another important consequence is gallstone pancreatitis. Occasional patients (usually elderly) present with painless progressive jaundice simulating a carcinoma of the pancreas. Some patients with bile duct calculi have no symptoms and present incidentally with abnormal liver function tests. There may be tenderness over the liver and pyrexia. The gallbladder is usually impalpable.

Investigations reveal cholestatic liver function tests and leucocytosis. Calcified gallstones may be seen on plain abdominal X-ray in 20 per cent of patients with cholelithiasis. An ultrasound examination (or CT) usually reveals a dilated CBD and calculus. MRI or endoscopic ultrasound (EUS) give more detail if needed. ERCP or PTC (Fig. J.3) are used for stone removal or stent insertion.

**Parasites**

The most important parasite is the roundworm *Ascaris lumbricoides*, which is a relatively common cause of obstructive jaundice in African children. It is released from an ovum in the duodenum and migrates into the intestinal wall, and hence the portal circulation. Worms enter the liver, heart and lungs, migrate into the pharynx, and are swallowed. Patients present with haemoptysis, bronchitis and pneumonia; occasionally, a worm blocks the bile duct to cause jaundice, cholangitis or act as a nidus for the development of a calculus.

**Causes affecting the wall of the bile duct**

**Bile duct trauma**

This is usually a consequence of inadvertent division at operation, (usually cholecystectomy).

**Cholangiocarcinoma**

These are epithelial tumours of the bile duct and arise at any point within the biliary tree. The intrahepatic type is increasing in incidence and can be difficult to distinguish from hepatocellular cancer. Patients present with pruritus, mild abdominal discomfort and obstructive jaundice. One important variant is a tumour arising at the bifurcation of the main left and right hepatic duct (Klatskin tumour). This tends to be slow-growing and carries a better prognosis than tumours arising elsewhere in the biliary tree.

**Carcinoma of the ampulla**

Tumours arising from the ampulla are uncommon and they present with painless obstructive jaundice. They are associated with familial adenomatous polyposis, are more slowly growing than carcinoma of the pancreas, and can be treated with a Whipple’s operation (pancreaticoduodenectomy).

**Biliary atresia**

This presents as deepening obstructive jaundice within 2–3 days of birth. Liver failure develops by the age of 3–6 months. The diagnosis is made by hepatobiliary imino-diacytic acid (HIDA) scanning, followed by cholangiography.

**Sclerosing cholangitis**

This forms part of the newly recognized IgG4-related diseases and is characterized by lymphoplasmocytic infiltration of the bile duct. Steroids are usually very effective in the short term.

*Figure J.3* Percutaneous transhepatic cholangiography. The bile ducts are dilated, and the filling defect caused by a stone can be seen at the lower end of the common bile duct. Multiple stones are also seen in the gallbladder.
Causes compressing the bile duct from outside
Carcinoma of the pancreas

The most important cause is carcinoma of the pancreas, which invades the common bile duct as it passes through the head of the gland. Cancers in the head of pancreas tend to present earlier than those in the body or tail as obstruction of the common bile duct and therefore jaundice is an early feature in the former. Patients present with weight loss, jaundice and some with progressive and continuous pain in the back, due to invasion of the coeliac plexus and other retroperitoneal structures.

Examination may reveal cachexia, jaundice and a palpable gallbladder (Fig. J.4; see also Fig. G.4). There may be evidence of metastatic spread; in particular, there may be supraclavicular lymphadenopathy (Virchow’s node) or tumour nodules in the umbilicus (Sister Joseph’s nodule).

Most tumours arise from ductular epithelium and carry a very poor prognosis. It is important, however, to remember that some tumours are more amenable to therapy; these include the cystadenocarcinoma, a tumour of relatively young women, and the apudoma. Ultrasonography, CT or MRI usually reveals a dilated common bile duct (and often pancreatic duct – ‘double duct sign’) and mass within the pancreatic head. ERCP reveals a stricture within the head of the pancreas, corresponding to a low bile duct malignant stenosis.

Malignant obstruction of the extrahepatic bile duct may also be due to enlarged lymph nodes in the region of the porta hepatis. Causes include breast, myeloma or intra-abdominal cancer or lymphoma.

Mirizzi’s syndrome

This is a rare entity in which a stone impacts in Hartmann’s pouch (gallbladder outlet), causing compression and obstruction of the common hepatic duct.

Cysts

Hydatid cyst is a rare cause of obstructive jaundice; far more commonly, these are found incidentally by plain X-ray. Simple cysts or choledochal cysts may also cause obstructive jaundice.

JAW, DEFORMITY OF

Leandros Vassiliou & Mark McGurk

Jaw deformity may arise from congenital or acquired conditions; for a discussion of the latter condition, (see JAW, SWELLING OF, p. 314). The reader should refer to this section for details of pathological conditions causing jaw deformity. Trauma to the jaws may cause deformity due to misalignment of the bone fragments. The majority of conditions to be considered here, however, are of a developmental nature occurring before birth or during the growth period.

Congenital conditions:

- Cleft palate
- Vascular malformations
- Pierre Robin syndrome
- Hemifacial microsomia (First arch syndrome)
- Treacher Collins syndrome
- Premature synostosis
- Achondroplasia
- Hemifacial atrophy

Acquired conditions:

- Disorders of the temporomandibular joint
- Acromegaly
- Cysts and odontogenic tumours (see JAW, SWELLING OF, p. 314)
- Radiation growth inhibition

The mandible forms in utero around a rod of cartilage known as Meckel’s cartilage. This is subsequently replaced by bone but with cartilaginous remnants persisting in the mandibular condyle. As with other growth centres, elongation of the mandible occurs by
the growth of this cartilage with secondary ossification that is completed by surface deposition of bone and remodelling of the bone unit.

The remaining facial skeleton is formed in fibrous bone, and growth occurs by a combination of sutural growth, surface deposition/remodelling, and cartilaginous growth with secondary ossification. The main stimulus for growth of the maxilla is the growth of the brain, causing an increase in size of the cranial vault and the cranial base to which the maxilla is joined. The cranial base also increases in length by cartilaginous growth at the sphenop-occipital synchondrosis. Growth within the maxilla is stimulated by the development of the nasal capsule and the eyes.

The inherited pattern of growth is complimented and moulded by functional forces provided by the eruption of teeth and muscles of mastication. A defect in one or more of these components will produce facial deformity. The commonest jaw deformity (if it can be described as such) arises from a simple imbalance between the growth of the maxilla and the mandible to produce variable degrees of dental malocclusion (Fig. J.5).

CONGENITAL JAW DEFORMITIES

Cleft palate

Cleft palate occurs in approximately one in every 700 live births and is due to a failure of growth and fusion of the palatal shelves in the embryo. Females are affected more than males, and there may be an associated cleft of the lip (Fig. J.6). There is a genetic disposition to this deformity, with a strong family history, but other exogenous factors have been implicated. These are drugs such as retinoids, phenytoin, methotrexate and other anti-folate agents. As folic acid deficiency is implicated with this abnormality, all mothers should be given folic acid supplements at conception. About 5 per cent of cases of cleft lip and palate are associated with other congenital abnormalities.

Vascular anomalies

Vascular lesions comprise a heterogeneous group of malformations that involve the arterial, capillary,
venous, lymphatic vessels or any combination. The classification is evolving but an accepted version is that of Waner and Suen. There are two main families: haemangiomas (subdivided into superficial, deep-seated and mixed type) and vascular malformations (capillary, arterial, venous and lymphatic or combinations). Haemangiomas are subdivided again into congenital and infantile. The former are split into rapid involuting and noninvoluting congenital haemangiomas, both of which are present at birth but the first involutes in the first few months of life while the other persists. In contrast, the infantile haemangioma is a true benign neoplasm that has three distinct phases of growth and is not present at birth. It has two rapid growth phases, one 1–2 months after birth, and the second at 4–5 months followed by an involution phase. Growth may be rapid and unpredictable. Haemangiomas respond to propanolol, and surgery is held in reserve especially for some rapidly proliferating lesion in aesthetic areas. This subject has moved out of the world of the generalist into specialist practice.

Vascular malformations are present at birth, and the capillary form (port wine stain) is amenable to laser therapy. The venous variety is present at birth and has no growth spurts but simply enlarges as the patient grows. They are amenable to laser and sclerosant therapy. Arteriovenous (AV) malformation can be in soft tissue, bone or both. Endovascular embolisation is the treatment of choice, with or without surgery. Simply ligating the feeding artery is a mistake and aggravates the situation.

Capillary malformations present as superficial ‘port wine’ stains and are associated with Sturge-Weber syndrome.

Lymphatic malformations (Fig. J.7) are low-flow and can be micro-cystic (previously known as lymphangiomas) or macro-cystic (cystic hygromas). They may enlarge with age and can induce hypertrophy of the tissues. Huge lesions can compromise the airway in infancy and dictate surgical debulking. Surgery/laser ablation has a part to play in controlling this disease but sclerotherapy is also effective.

High-flow malformations such as arteriovenous malformations (AVMs) present as pulsatile masses. Endostable growth may cause jaw deformity. They may bleed spontaneously with catastrophic consequences. Schobinger has described four clinical stages based on the expansion and the complications of AVMs: Quiescence, Expansion, Destruction (when the AVM may spontaneously bleed or cause tissue necrosis) and Decompensation (when cardiac failure may occur as a result of haemodynamic changes due to the AVM). Complex malformations may be regional or diffuse and are usually part of a syndrome (von Hippel–Lindau, Kipple–Trenaunay, Maffucci, Parkes Weber).

Pierre Robin syndrome
This syndrome is thought to be caused by hypoplasia of the mandible, preventing the normal descent of the tongue and thus preventing fusion of the embryonic palatal shelves. The syndrome is therefore characterized by a small mandible, cleft palate and protruding tongue. The baby may present with feeding and respiratory problems, which can be corrected by the construction of a small dental plate and nursing in the supine position.

Normal growth of the mandible can be anticipated after a few years, once any palatal defect has been closed.

Hemifacial microsomia
Hemifacial microsomia is also known as Goldenhar syndrome. It is associated with abnormal development of the first and second branchial arches and characteristically exhibits a deformed or absent ear, macrostomia and an underdevelopment of the mandibular ramus and condyle. The masticatory muscles on that side are also deficient, and there is hypoplasia of the orbit and zygoma on the ipsilateral side (Fig. J.8). Other associated abnormalities may be present, particularly of the vertebrae. The long held mechanism of pathogenesis through a stapedial artery bleed in utero has been superseded by a genetic based explanation.
Treacher Collins syndrome
This is an inherited autosomal dominant condition affecting the facial skeleton in a similar way to the first arch syndrome, but the abnormalities are bilateral and symmetrical. Due to the poor development of the zygomatic arches, prominent nose and small jaw, many of the patients have a fish-like appearance.

Premature synostosis
Premature fusion of the cranial sutures is a feature of Crouzon's and Apert's syndromes. These are autosomal dominant conditions associated with mutations of the FGFR2 gene. The associated lack of growth of the cranial base as a result of synostosis leads to extreme underdevelopment of the mid-third of the face and various degrees of exorbitism and hypertelorism respectively.

Achondroplasia
This rare condition usually represents a sporadic mutation of the FGFR3 gene; less than 20 per cent will be of a familial nature. The aetiology is not completely understood, but there is a defect of endochondrial ossification. Failure of growth at the spheneno-occipital synchondrosis and lack of growth in the maxilla produce the characteristic underdevelopment of the mid-third of the face. Curiously, growth of the mandible is unaffected, leading to relative mandibular prognathism.

Hemifacial atrophy
A peculiar form of facial asymmetry (Parry–Romberg syndrome) arises with slow progressive atrophy of the soft tissues of one side of the face (coup de sarbe) with secondary deformity of the facial skeleton. Patients may also exhibit contralateral Jacksonian epilepsy and trigeminal neuralgia. This condition is thought to be due to an abnormality of the sympathetic system and is often associated with scleroderma.

ACQUIRED JAW DEFORMITIES
Disorders of the temporomandibular joint
The mandibular condyle may be affected by trauma, infection from the middle ear or juvenile arthritis, all of which can adversely affect the condylar growth centre. The result is underdevelopment of one side of the mandible with compensatory growth on the contralateral side with the result the chin point moves towards the damaged condyle. This produces a facial asymmetry and underdevelopment of the ipsilateral side of the face in the vertical plane and secondary changes in the maxilla. Fractures of the temporomandibular joint may, on occasions, be followed by ankylosis, and this, too, will prevent normal development of the affected side of the face (Fig. J.9). Treatment of all these conditions is by surgical correction.

In some patients, there may be excessive growth of the condyle, known as condylar hyperplasia, resulting in asymmetry of the facial skeleton, but this time with the chin point pushed to the side of the normal condyle.

Figure J.8 First arch syndrome, classified by the severity of failed development of the mandible; however, growth defects also occur in the zygoma and temporal bone.
Acromegaly follows autonomous hypersecretion of growth hormone caused by hyperplasia or an adenoma of the pituitary acidophil cells. The face is invariably affected by this condition, with overgrowth of the mandible to produce prognathism with malocclusion, enlargement of the tongue and deposition of bone at the supraorbital ridges and zygomas.

The facial skin also becomes thickened, as does the subcutaneous tissue, producing an accentuation of the normal skin folds. The nose becomes enlarged, especially at the tip, as do the lips. Treatment should be for the underlying pituitary problem, and corrective jaw surgery should only be undertaken following stabilization of the condition. Many of these patients have a cardiomyopathy, and there may be serious complications during anaesthesia.

**Radiation growth inhibition**

Ionizing radiation is used as part of multimodality treatment for paediatric head and neck malignancies. In the growing craniofacial skeleton, radiotherapy causes growth inhibition and subsequent deformity in over 66 per cent of long-term survivors.

The skull is basically divided into neurocranium (mainly the vault, forming a bony case around the brain) and viscerocranium, which is the skeleton of the face. The neurocranium is further divided into ‘membranous neurocranium’ consisting of the flat bones around the brain and ‘chondrocranium’, which essentially constitutes the skull base.

The membranous neurocranium arises by intramembranous ossification, the chondrocranium by endochondral ossification and the viscerocranium by intramembranous ossification. By 5–7 years of age the brain stops expanding and the child reaches its cranial capacity. The viscerocranium continues to develop and the face grows with subperiosteal surface bone deposition and internal resorption. This process accommodates the growth of the paranasal sinuses and the dentition, and it continues until adulthood.

The differential in the neurocranium and viscerocranium growth and ossification timings results in the upper face (which consists mainly of the frontal bone – neurocranium) completing its growth by the age of 5–7 years, while the rest of the face (middle and lower third – viscerocranium) continues to develop until the age of 16–18 years.

When it is employed to treat a head and neck malignancy, radiotherapy has a profound effect on the development of the craniofacial skeleton. Radiation diminishes vascularity, decreases cellularity of all bone cell populations and suppresses osteoblastic and osteoclastic activity. In addition, cranial radiation may cause growth hormone deficiency leading to generalized growth retardation.

The overall pattern of growth inhibition may range from dental abnormalities to hemifacial or panfacial microsomia, depending on the site of irradiation, the age when radiotherapy is applied and the radiation...
protocol. In general, the abnormalities caused are greater in severity the earlier the age of treatment and the higher the radiation doses.

**MANAGEMENT OF JAW DEFORMITY**

The facial appearance and jaw function of patients with jaw deformity can be improved by combined orthodontic and surgical correction. Minor deformity can be corrected by movement through osteotomy procedures, while the more severe cases with significant abnormalities of the soft tissues should be treated by distraction osteogenesis techniques or complex craniofacial surgical procedures.

**JAW, PAIN IN**

Leandros Vassiliou & Mark McGurk

Pain in the jaw mostly arises from the dental structures and their supporting bone, the temporomandibular joint and the associated muscles of mastication. In the upper jaw, infections of the nose and paranasal sinuses may additionally cause midfacial pain. Disorders of the trigeminal nerve are a relatively rare cause of facial pain, but atypical facial pain, which is part of a psychological illness (usually depression), is quite common.

**DENTAL PAIN**

Inflammation in the pulp chamber of a tooth can be caused by dental caries, inadequately insulated restorations or iatrogenic trauma from a drill. Pressure within the chamber rises, threatening a compartment syndrome. The tooth becomes hypersensitive to thermal stimulation and, if necrosis commences, the pain becomes intense but with the characteristic feature that it is poorly localized. It is difficult to identify the offending tooth.

After a number of days, the pain abates and the tooth then becomes tender to palpation as the inflammatory proteins pour out of the apex to sensitize the suspending periodontal ligament. Symptoms are more in keeping with a low-grade infection, where the pain is moderate to severe, throbbing in nature and easily localized (Fig. J.10) (see JAW, SWELLING OF, p. 314).

Periodontal abscess or pericoronal infection around a partly erupted wisdom tooth has a similar pattern of symptoms which is quite different from odontogenic pain.

Developmental lesions within the jaw (e.g. odontogenic cysts; see JAW, SWELLING OF, p. 314) may become infected. The diagnosis is usually readily apparent following a careful history followed by clinical and radiographic examination. Should the abscess involve the masticatory muscles and especially the masseter, then trismus will be present together with signs of acute infection.

**ACUTE POST-EXTRACTION OSTEITIS**

The pain from ‘dry socket’ is excruciating in nature. It commences 2–3 days after the extraction of a tooth, and is due to an osteitis of the cortical bone lining the socket, which is the origin of the pain. The blood clot cannot attach to the bone and is lost, hence the name dry socket. Patients look ill but instantaneous relief can be obtained by packing the socket with a dressing containing oil of cloves. The socket should then be irrigated to remove debris and the oil of cloves dressing repeated for 2 or 3 days. The condition is self-limiting but healing is delayed for weeks due to the effect of the oil. Antibiotics are not normally required.

**BONE PATHOLOGICAL CONDITIONS**

Powerful pain receptors line the periosteum and, when the membrane is disrupted, an intense then a deep boring pain ensues. Injury can be of a surgical nature or caused by trauma, especially when bone fragments are mobile. Reduction (open or closed) and immobilization of the bone fragments are the treatment of choice.

Pathological fractures secondary to osteolytic lesions, osteoradionecrosis, primary malignant or metastatic disease present with similar symptoms.

Sickle-cell anaemia is a hereditary condition caused by a point mutation in the gene encoding the β-globin chain of haemoglobin, leading to the production of the abnormal haemoglobin S (HbS). Under specific conditions, such as hypoxia, dehydration, acidosis or infection, haemoglobin S changes conformation, distorting the erythrocytes into a sickle shape.
As a consequence, the blood cells lose their elasticity and, becoming rigid, cannot pass through narrow capillaries. This precipitates capillary obstruction and a vaso-occlusive crisis which is acutely painful. Sickle-cell crisis can affect the jaws, especially the mandible, causing painful swelling. As a result of the ischemia, inferior dental nerve paresthesia or even avascular necrosis of the bone may occur. The treatment is usually symptomatic with hydration, analgesia, blood transfusions and addressing the precipitating factor of the sickle-cell crisis.

**TEMPOROMANDIBULAR JOINT**

Dislocation of the temporomandibular joint causes jaw pain which persists until the joint is reduced either under local anaesthetic or, more commonly, under sedation or even general anaesthetic.

The temporomandibular joint may be affected by any of the conditions that afflict the other joints, for example rheumatoid arthritis, osteoarthritis and septic arthritis. In this case, pain and swelling in the acute phase are present in the preauricular region, with limitation of jaw movements. An acute effusion in the temporomandibular joint may arise as a result of a blow to the jaw or a fracture involving the joint. This again will cause pain and swelling anterior to the ear.

The most common cause of pain at this site is temporomandibular joint dysfunction syndrome. This is an extremely common and much written about condition, without the aetiology and pathogenesis of the condition being clearly understood. It is multifactorial in origin and the contribution of the various factors varies with the individual. The physical effect can be loosely described as a strained joint with ligaments stretched, insertions torn, inflammatory effusion lowering receptor pain threshold and condylar disc hypermobility.

Stress and psychological mindset are contributory factors that lead to muscle tension and parafunctional habits (bruxism). The muscles of mastication are contracted for long periods leading to spasm and pain. Fibres of the lateral pterygoid muscle are inserted into the meniscus of the temporomandibular joint and, during teeth clenching, the muscle fires, gradually stretching the ligaments and ultimately pulling the disc off the head of the condyle if the habit is not contained. Loose discs produce a click on movement of the jaw, and further displacement causes locking of the joint with significantly reduced mouth opening. Acute events, such as a blow to the jaw or a road traffic accident where the car is struck from behind, can cause hyper-extension and initiation of symptoms that are then sustained by other factors, such as an altered occlusion, periods of wide mouth opening or chewy foods. None of these is a primary initiator but it can be a secondary factor. Pain symptoms fluctuate over time. Pain may originate from the temporalsis muscle and radiate behind the eye where it is interpreted as a headache. Pain is also experienced within the ear and in the preauricular region. It is not uncommon for an audiology opinion to be erroneously sort for these symptoms. Tenderness may be elicited intraorally at the upper insertion on the masseter muscle and at the anterior margin of the medial pterygoid muscle. Prolonged contraction of the muscles may induce muscular hypertrophy, especially of the masseter muscle, which produces a characteristic swelling at the angle of the jaw.

The natural history of this condition is for it to resolve over time. It is a disorder of young adults and is mostly passed by age 50 years. The best results come when the patient has insight into the origin of the condition and works with the clinician to minimise aggravating factors. The treatment usually consists of an in-depth explanation of the condition, the use of an occlusal bite guard and NSAIDs. Interventional approaches should be conservative in nature and reversible (e.g. arthrocentesis or arthroscopy of the joint). Surgical exploration of the joint should be left to the expert who has specialized in this joint. Surgical intervention can be fraught with problems including facial palsy and a proclivity to medico-legal complaints.

**ACUTE MAXILLARY SINUSITIS**

Pain arising in the acutely infected maxillary sinus may be confused with pain of dental origin as the tooth roots of the upper teeth have a very close relationship to this structure. Maxillary sinusitis normally follows an upper respiratory tract infection, especially if the normal drainage of the antrum through the ostium is reduced. However, maxillary sinusitis can arise from dental infection if the root and abscess point into the maxillary antrum rather than into the oral cavity. Infection of the maxillary sinus may also follow the creation of an oro-antral fistula after dental extraction.

In maxillary sinusitis, the patient suffers from a throbbing pain in the cheek, which is aggravated by bending down, but there is hardly ever associated facial swelling, only occasionally erythema of the conjunctiva. The teeth may be tender to percussion. Intranasal examination may demonstrate a mucopurulent discharge through the normal ostium. The healthy maxillary sinuses are translucent on X-ray examination. Opacification is indicative of disease. Infective changes are invariably bilateral and consequently unilateral opacification demands investigation in case an occult tumour is present (Fig. J.11).
A neoplastic process of the maxillary antrum should be suspected if the pain does not respond to normal measures, or if there is swelling of the face or oral cavity associated with bone destruction and displacement of the teeth. A tumour of the maxillary antrum may also cause epistaxis and sensory loss in the maxillary division of the trigeminal nerve. If these signs are present, a biopsy should be obtained either endoscopically through the nose or via a Caldwell–Luc approach.

PRIMARY NEURALGIAS
Primary neuralgias may be defined as the disturbed function of a nerve without the existence of any recognized aetiological factor or pathological process along the nerve pathway or its central connections. In the jaws, the trigeminal nerve is affected, and very occasionally also the glossopharyngeal nerve. There are no associated signs, and the diagnosis is made from the history.

The pain characteristically affects only one branch of the trigeminal nerve initially, although later in the disease it may spread to two or occasionally three divisions. The disease most commonly affects patients aged over 60 years, and the incidence is twice as common in women as in men. The diagnosis should not be accepted in those under 50 years of age without extensive investigation.

The pain is very severe and is described as sharp, or similar to an electric shock. It is paroxysmal in nature, lasting for only a few seconds with intervals of a few minutes or a few hours. The pain may be felt spontaneously or in response to stimulation within a specific area (trigger zone) on the face. This trigger zone may be activated by a cold wind, shaving, washing, eating or cleaning the teeth. Natural remission is fairly common.

The main differential diagnosis is between causes of dental pain, and these should be excluded before a diagnosis of trigeminal neuralgia is made. Treatment is with drugs, such as carbamazepine, epanutin or gabapentin. Response to these drugs assists in the diagnosis. The drug carbamazepine occasionally causes agranulocytosis, and therefore regular monitoring of the drug plasma levels as well as the white blood cell count is essential. In every case, a careful examination of the central nervous system should be made, which often includes a magnetic resonance imaging (MRI) scan to exclude a neoplasm (neuroma) or pulsatile intracranial aneurysm as the cause. In the younger age group, multiple sclerosis needs to be excluded as it can mimic trigeminal neuralgia in the early stages of the disease. Should medical treatment fail, a neurosurgical approach can be considered. In contrast, the results of peripheral nerve ablation are unsatisfactory and unpredictable.

SECONDARY NEURALGIAS
Here, an identifiable pathological process is acting at some point along the trigeminal nerve or its central connections, producing pain at the periphery. The symptoms may be similar to trigeminal neuralgia, or it may be a duller, more continuous pain. Neoplasms are the most significant cause, other examples being aneurysms or compression of the nerve in the bony canal in Paget's disease.

Accurate examination of the cranial nerves is essential, followed by a full clinical and radiographical examination, including computed tomography or MRI scans to establish the diagnosis.

POST-HERPETIC NEURALGIA
Involvement of one of the branches of the trigeminal nerve with the herpes zoster virus will produce pain and vesication in the anatomical distribution of that nerve. Once this attack has resolved, scarring of the involved nerve may leave the patient with post-herpetic neuralgia and possibly sensory disturbance. This pain can be severe and resistant to treatment, but neuropathic drugs are the treatment of choice and should be introduced early to stop pain becoming established.
MIGRAINE
Migraine and migrainous neuralgia may occasionally involve the upper jaw, although the predominant features are manifested as headache. The diagnosis is normally made from the history when an intense pain is associated with visual disturbance, nausea and constitutional symptoms. In migrainous neuralgia, the pain is predominantly behind the eye, and patients may also experience pain in the maxilla and temple regions. There may be watering of the eye and flushing of the facial skin.

REFERRED PAIN
There are two important examples of referred pain to the jaws. One is that of myocardial ischemia due coronary artery insufficiency, which may produce pain in the left side of the mandible the other is referred pain to the ear from invasive lesions in the tongue and pharynx. An elderly patient who complains of a sore throat and ipsilateral earache (otalgia) has cancer until proven otherwise.

ATYPICAL FACIAL PAIN
A relatively small but well-established cohort of patients present with atypical facial pain, and many have an unusual psychological profile. The pain is described as being very severe, but it does not produce any restriction upon the normal function of the jaws and oral cavity. It does not have an anatomical distribution, commonly involves both sides of the face and jaws, and moves from one part of the facial skeleton to another. Typically the pain seems to arise from a sound tooth. Despite treatment, the pain persists and the tooth is removed. The pain then moves to the next tooth and after two or three extraction it dawns on the dentist that the pain is not of dental origin. The pain does not respond to analgesics, and usually there are many other associated symptoms such as a dry mouth, burning tongue, and other complaints throughout the body.

There can be some overlap between this condition and temporomandibular joint dysfunction caused by stress, but, in every case, it is essential to exclude pain due to any one of the other causes just described. Therefore, atypical facial pain is often a diagnosis of exclusion and, once made, any underlying depression should be treated by medication or referral for psychiatric help. (See also FACE, PAIN IN, p. 182.)

JAW, SWELLING OF
Leandros Vassiliou & Mark McGurk

Swellings of the jaw, once they have reached a certain size, will be obvious as facial swelling. The true nature will be ascertained by taking history, performing an intra- and extra-oral examination and obtaining radiographs. Smaller swellings may be visible only on examination of the oral cavity or will have been discovered by the patient. Testing of the trigeminal cranial nerve should always be carried out since a change in sensation may have very significant implications. Swellings of the jaw can be incorrectly diagnosed by the inexperienced clinician as swellings of the submandibular salivary gland or submandibular lymph nodes or, in particular, as a parotid mass.

INFECTION ASSOCIATED WITH THE DENTAL STRUCTURES
Bacterial infection associated with the dental structures is by far the most common cause of swellings of the jaw. An alveolar abscess arises when gangrene of the dental pulp occurs following dental caries, extensive dental restorations or trauma (Fig. J.12). This infection then spreads to the alveolar bone to cause a localized osteitis but remarkably, in the majority of cases, does not cause osteomyelitis. Instead, the abscess, as it enlarges, becomes localized and perforates either the lateral or medial plate of the outer compact alveolar bone. It is at this stage that it presents as a swelling of the jaw, which is tender and covered by inflamed mucosa. Occasionally, an alveolar abscess is associated with sensory loss of the mandibular branch of the trigeminal nerve.

A periodontal abscess arises from bacterial infection within the periodontal membrane of the tooth, which is usually associated with previous chronic periodontal disease. A pericoronal abscess arises in the mucous membrane surrounding the crown of an erupting or impacted tooth, the majority being associated with wisdom teeth. At this stage, the swelling is largely confined to the region of the jaws and may

Figure J.12 Intraoral dental abscess in the palate.
discharge intraorally. However, should the bacteria gain access to the adjacent soft-tissue compartments, facial cellulites or soft-tissue abscess formation will follow (Fig. J.13). Depending upon the anatomical position of the infection, the submandibular area may become swollen, or the cheek (buccal space) or more posteriorly the submasseteric space, which may be misdiagnosed as a parotid swelling.

On the medial aspect of the jaw, swelling may occur in the sublingual space or more posteriorly in the pterygoid, lateral pharyngeal or peritonsillar space. Diagnosis of these latter space infections may be difficult and, if beneath a muscle compartment, are associated with severe trismus.

An orthopantomogram X-ray is required in all cases of facial swelling to establish or eliminate a dental cause. The majority of swellings will respond to antibiotic treatment of an underlying dental cause and surgical drainage where necessary. The medial swellings are potentially very serious, as airway obstruction may follow unless the neck is decompressed in severe cases. Ludwig’s angina is an acute emergency in which bilateral sublingual and submandibular and submental spaces are involved in acute infection. Again, urgent surgical decompression of the neck is required to prevent respiratory obstruction. In advanced cases, a surgical tracheostomy may be the only way to secure the airway.

Persistent recurrent infection causing chronic swelling and discharge may be due to actinomycosis, which is an opportunistic infection.

True osteomyelitis of the jaws is now relatively rare following the improvement in general dental health and the use of antibiotics. The overarching feature in the early lesion is intense and intractable pain (like a dry socket which is a similar phenomenon) and constitutional changes (tachycardia, temperature, raised white cell count), but little to see in the mouth. When established, the pain is accompanied by loosening of the adjacent teeth and usually there is sensory compromise (paresthesia) of the respective branch of the trigeminal nerve (inferior dental nerve). In time (weeks), the overlying mucosa becomes swollen and hyperaemic, and, indeed, sinuses may develop through which there is a discharge of pus and bony sequestra. In the acute phase, it is invariably associated with significant soft-tissue swelling. The radiographic changes take some time to develop but show diffuse rarefaction and sequestrum formation. Over a period of months, subperiosteal woven bone may become a feature and the infected bone may be subjected to a pathological fracture.

Management of all these infections is by high-dose antibiotics in the acute phase with elimination of the source of infection. In established lesions, the sequestra have to be removed and the defects restored. In ideal circumstances, there are new options such as hyperbaric oxygen to stabilize the wound and grafting the site with new vascularized bone.

TRAUMA
Fractures of the mandible are relatively common and are associated with swelling caused by haematoma formation from the bleeding marrow spaces and the peristomeum. The swelling may be made worse by a protruding bone fragment or the presence of a foreign body. The injury may be insufficient to cause a fracture, but it may nevertheless produce a haematoma in the soft tissue. This normally resolves without treatment but, occasionally, may require aspiration or drainage. The diagnosis of a fracture is normally easy to make from the history, the abnormal mobility of the fragments and the irregularity of the dental arches. In many fractures, a laceration of the oral mucosa is also present. The diagnosis is confirmed by radiographic examination (Fig. J.14).

CYSTS
Cystic lesions are very common in the jaws and are an important cause of jaw swelling. These are classified as developmental or inflammatory.

An inflammatory cyst develops in association with a dental or periodontal infection generating an inflammatory process in the periapical tissues or the periodontium. These cysts are termed radicular or periodontal, respectively. As these cysts arise from odontogenic cell remnants in the bone stimulated
by the inflammation and progressively enlarge, the alveolus expands and, just before the bone perforates, the phenomenon of 'eggshell cracking' can be elicited and the swelling is naturally fluctuant. The cyst may slowly enlarge, or, should it become infected, acute swelling with inflammation occurs.

The developmental cysts are subdivided into odontogenic or non-odontogenic, depending on the origin of the epithelium lining the cyst. Jaw cysts of odontogenic origin are the dentigerous cyst (follicular), the lateral periodontal cyst, the eruption cyst, the gingival cyst and the glandular odontogenic cyst. Dentigerous cysts are particularly common, always associated with an unerupted tooth. They arise from the natural follicle that covers the crown of the tooth and helps it to erupt. The follicle enlarges but always with a tooth attached, which is the diagnostic feature of this cyst (Fig. J.14). The cyst slowly enlarges and may cause displacement of the involved tooth or the adjacent teeth.

A common but rather problematic jaw cyst due to its high rate of recurrence was the previously named odontogenic keratocyst. As a result of its biological behaviour and proclivity for recurrence, it has now been reclassified as a benign tumour, and it is termed a keratocystic odontogenic tumour.

Non-odontogenic developmental cysts include the cyst of the nasopalatine duct and the nasolabial cyst which are frequently encountered in the maxillofacial region (Fig. J.15).

SWELLING DUE TO ODONTOGENIC CAUSES/TUMOURS

Unerupted impacted teeth are a frequent cause of jaw swelling; these are commonly supernumerary teeth or the canine and premolar teeth in the palate and the premolars in the lower jaw. In the elderly, when the molar teeth have been lost, an unerupted wisdom tooth may produce a swelling in the posterior area of the alveolus.

A long list of odontogenic tumours can cause jaw swelling. The classification according to WHO divides them into benign and malignant with the former being most common. The histological classification is based on the primary odontogenic tissue of origin and its grade of maturation. Benign odontogenic tumours include ameloblastomas, adenomatoid odontogenic tumours, keratocystic odontogenic tumours (previously known as odontogenic keratocysts), ameloblastic fibromas, dentinogenic ghost cell tumours, odontogenic myxomas and cementoblastomas, and a number of malignant odontogenic tumours such as the ameloblastic carcinoma, the clear cell odontogenic carcinoma and the ghost cell odontogenic carcinoma that are extremely rare. In the following paragraphs the most common benign odontogenic tumours will be described.

Odontoma

*Odontomas* are hamartomas of odontogenic origin and may give rise to a swelling of the jaw. The diagnosis is confirmed by X-ray. Odontomas can be compound when displaying similar tissue complexity as a normal tooth but random shape, or complex when they present as an amorphous conglomerate of dental tissues.

Ameloblastoma

Ameloblastoma is a benign neoplasm of the odontogenic epithelium (ameloblasts) and is locally invasive. Unless infection supervenes, the tumour is quite painless as it slowly enlarges. In developing countries, the lesions may reach enormous proportions before assistance is sought. Radiographs usually show a multilocular radiolucency, but unilocular variants
may present. Spacing of the teeth may occur, but sensory disturbance of the trigeminal nerve is not usually a feature. As these tumours tend to recur, treatment is by resection with a margin of healthy bone and bone graft reconstruction.

**Odontogenic myxoma**
Another benign tumour to affect the jaws and cause expansion is the myxoma, which is a rare benign neoplasm arising from odontogenic mesenchyme. It produces a multilocular radiolucent appearance with multiple criss-crossing septi in the defect. Treatment is by surgical excision.

**SWELLING DUE TO NON-ODONTOGENIC PATHOLOGY**

**Fibrous dysplasia**
This condition of bone of unknown origin is characterized by replacement with fibrous tissue and enlargement in all three dimensions. At this stage, the abnormal bone is very vascular, but subsequently ossification occurs to produce an amorphous radiopaque appearance on the radiograph. The process tends to cease at skeletal maturity. The jaws are frequently affected in monostotic fibrous dysplasia and may also be involved in polyostotic fibrous dysplasia and Albright's syndrome. This facial deformity is usually corrected by surgical sculpturing.

**Cherubism (familial fibrous dysplasia)**
Symmetrical enlargement of the facial skeleton occurs in this inherited condition, which is usually apparent in early life and then arrests at puberty. Radiographs show symmetrical multilocular radiolucent areas of the jaws and, histologically, the bone is replaced by fibrous tissue with multinucleated giant cells as a predominant feature. The giant cells, in some cases, make the differential diagnosis difficult from giant-cell granuloma or hyperparathyroidism. The blood chemistry is, however, usually normal.

**Paget's disease**
This disease of bone of unknown aetiology found in patients in middle to late life may affect the mandible but, more commonly, it affects the maxilla. According to some studies, approximately 15 per cent of cases show involvement of the facial skeleton. Enlargement of the facial bones may produce the characteristic 'leonine facies' and, intraorally, expansion of the dental alveolus occurs with displacement of the teeth. Pain may occur due to entrapment of the trigeminal nerve, and the radiograph shows the typical areas of patchy sclerosis. Extraction of the teeth can be difficult due to hypercementosis, and post-extraction bleeding may be severe. In the active phase of the disease, the serum alkaline phosphatase is raised, which will aid the diagnosis of the condition. Sarcomatous change in long-standing Paget's disease occurs but is relatively rare.

**Osteoma**
Osteomas of the jaw present as hard, round, bony swellings that may be endosteal (central) or subperiosteal (peripheral). Multiple osteomas of the facial bones are found in Gardner's syndrome, the other main feature being polyps of the large intestine, which have a tendency to become malignant. Radiologically, osteomas may be composed of dense radio-opaque bone or may have a high cancellous component, in which case they are relatively radiolucent. Torus palatinus is a developmental abnormality of the midline of the hard palate characterized by a cylindrical enlargement in the region of the midline palatal suture (Fig. J.16). Torus mandibularis is a similar, slowly enlarging developmental abnormality, but it arises on the lingual aspect of the mandible in the premolar region. All these osteomas are simply removed if they prove troublesome as a result of trauma to the overlying mucosa.

**Ossifying fibroma**
This is a benign fibro-osseous lesion, which causes a well-circumscribed mass of fibrous tissue showing areas of speckled calcification. It normally arises within the substance of the bone and slowly expands in all directions to produce a sclerotic margin. With time, the lesion becomes more calcified, and it can cause loosening of the adjacent teeth. Treatment is by surgical excision.

*Figure J.16* Torus palatinus.
**Cementifying fibroma**

This condition is similar in some ways to ossifying fibroma in that an area of bone is replaced by fibrous tissue, but subsequent calcification resembles dental cementum. Periapical cemental dysplasia is similar to cementifying fibroma but produces multiple sites of ossification with cementum. The diagnosis is usually easy to make from the radiographic appearances but it may, in some patients, produce a bony hard irregularity of the dental alveolus. Surgical removal is rarely required.

**Giant-cell granuloma**

The aetiology of this lesion is again unknown. It arises in the young age group and is confined to the tooth-bearing areas of the jaws, with the mandible as the most frequent location. The lesion can grow rapidly. The central giant-cell granuloma develops within the jaw, later perforating the alveolar bone to present in the mouth as a spherical purple swelling. Radiographs show a radiolucent area and, histologically, the tissue is vascular fibrous tissue with multinucleated giant cells. Confusion can arise with the brown tumour of hyperparathyroidism, but here the occurrence in the older age group with raised serum calcium and parathormone levels would indicate the diagnosis. Additionally, radiographs in hyperparathyroidism show multiple osteolytic lesions – osteitis fibrosa cystica – on skeletal survey. The peripheral giant-cell granuloma is described elsewhere (see GUMS, HYPERPARATHYROIDISM). The traditional treatment of both types is surgical excision but there are reports of the central giant-cell lesion responding to calcitonin or interferon therapy. This therapeutic approach can be considered in adolescents or young adults when the social consequences of a jaw resection must be taken into consideration.

**Vascular anomalies**

Endosteal vascular anomalies are more common in the mandible than the maxilla and, if truly endosteal, present as hard, non-tender, painless swellings (see also JAW, DEFORMITY OF). Haemorrhage may occur around the necks of the teeth, which may become loosened, and severe haemorrhage follows dental extraction.

**MINOR SALIVARY GLANDS PATHOLOGY**

Many hundreds of minor salivary glands are distributed widely throughout the oral mucosa, but they are found in abundance at the junction of the hard and soft palate. The important benign causes of enlargement are mucous extravasation cysts (confined to loose tissue) and tumours such as pleomorphic adenomas (Fig. J.17). Mucous extravasation cysts commonly occur in the lower lip and floor of the mouth.

**MALIGNANT TUMOURS**

Sarcomas of the jaw are rare, with the mandible being affected more often than the maxilla. The topic is now a complex area of study. The classification is divided into families with subgroups based on genetic criteria. Clinically low-grade tumours are difficult to distinguish from fibrous dysplasia or giant-cell tumours of bone. Normally, surgery is the treatment of choice in the indolent lesion. Rapidly enlarging swellings are high-grade lesions and are associated with drifting and loosening of the teeth and sensory disturbance of the involved trigeminal nerve. Radiographs demonstrate irregular destruction of the jaw, but in the osteogenic sarcoma there are, in addition, radiating trabeculae of new bone formation to give the characteristic ‘sun-ray’ appearance.

Intraosseous squamous cell carcinoma is rare but, when present, it causes a destructive lesion within the mandible on X-ray, sensory disturbance of the trigeminal nerve and expansion of the bone with loosening of the teeth (see also GUMS, RETRACTION OF). The majority of squamous cell carcinomas arise in the oral mucous membrane and produce an ulcer in conjunction with swelling of the soft tissue overlying the jaw, usually with underlying bone destruction (Fig. J.18).

An extra-nodal lymphoma presenting in the oral cavity is an uncommon event, but it should be suspected in an adult presenting with a painless, nondescript soft swelling that does not respond to simple measures.
(see also GUMS, HYPERTROPHY OF, p. 229). The soft tissues of the cheek, palate, gingivae and fauces can all be affected, without the regional lymph nodes being enlarged. The lesion has a tendency to grow quickly and secondary ulceration of the lesion can occur. A biopsy resolves the diagnostic dilemma. The incidence of oral lymphoma is significantly raised in patients with HIV infection. The adult lymphoma needs to be distinguished from Burkitt's lymphoma, which is commonly found in children in East Africa and is associated with the Epstein–Barr virus. Involvement of the jaw is the predominant site, with secondary spread into the adjacent tissues. This tumour responds well to chemotherapy.

The Kaposi sarcoma was relatively uncommon in the oral cavity until the advent of AIDS and HIV infection. This is a vascular tumour with a nodular, purplish appearance. The palate is the most frequent site in the oral cavity. It is particularly associated with homosexual males with HIV infection, and in this group it has become almost pathognomonic of AIDS, unless the individual is immunosuppressed for another reason. The lesion is thought to be caused by a herpes virus. Treatment is now medical in nature supported by radiotherapy.

Carcinomas of the maxillary antrum present late as they grow undetected until they burst out of the bony box of the antrum. The mode of presentation depends on the direction of growth: if downward, then a palatal swelling; if up, then proptosis; if anterior, then cheek swelling and numbness of the infra orbital nerve; if posterior, then trismus as ptyregoid muscles are invaded. Malignant minor salivary glands tumours (acinic cell, polymorphus low adenocystic, adenocystic and mucoepidermoid carcinoma) are normally slow growing and usually present as a discreet lump in the soft tissues or more commonly at the junction of the hard and soft palate. The diagnosis is confirmed by biopsy, which is ideally obtained by a dermatological punch which prevents tumour contamination of the adjacent tissues.

METASTATIC TUMOURS OF THE JAWS
Secondary deposits of tumours rarely affect the jaws, with the mandible as the more common site. Tumours of the bronchus, breast (Fig. J.19), kidney and thyroid produce osteolytic lesions, while carcinoma of the prostate tends to form an osteosclerotic deposit. The clinical signs are similar to other intrabony malignant tumours with swelling, pain, loosening of the teeth and sensory disturbance. Curiously, in the mandible the tumour, emboli deposit themselves at the entrance to and exit of the inferior dental canal. Over time a pathological fracture may occur.

Secondary deposits may also develop in the overlying soft tissue of the jaws, especially at the gingival margin, and these may be confused with a benign epulis. Rapid growth usually occurs to produce a fleshy mass, which may be friable and haemorrhagic. These lesions should be excised and submitted for histological examination lest the diagnosis be missed.

JOINTS, AFFECTIONS OF
Toby Garrood
Joint affections may be acute, as in rheumatic fever, gout or traumatic hydrarthrosis; acute relapsing becoming chronic, as may be seen in rheumatoid arthritis or reactive arthritis; chronic with an acute onset, as in some cases of generalized osteoarthritis; or insidious and chronic, as in most cases of osteoarthritis. The affection may be of one joint, a monoarthritis, as may be seen in tuberculous arthritis; or it may be the start of a polyarthritis, as is not infrequently seen in psoriatic arthropathy, in which a number of joints are eventually affected. The pattern
of joint involvement may be important: rheumatoid arthritis affects initially and chiefly the peripheral joints; ankylosing spondylitis affects the sacroiliac joints and spine with or without peripheral joint involvement; and polymyalgia rheumatica affects the pelvic and shoulder region. The terminal interphalangeal joints are affected commonly in osteoarthritis (Heberden’s nodes) and psoriatic arthropathy, but rarely in adult rheumatoid arthritis. Interphalangeal involvement of the toes is unusual in rheumatoid arthritis, but more common in seronegative arthritis. Joint involvement may be in the form of a flitting and transient polyarthritis, as in rheumatic fever, some cases of systemic lupus erythematosus and a number of other conditions, or in a recurring or palindromic pattern subsiding completely between episodes, as in some cases of rheumatoid arthritis. Joints may be swollen because of:

- Bony enlargement, as in osteoarthritis or Charcot’s joints in tabes dorsalis or syringomyelia
- New periosteal bone deposition, as in hypertrophic pulmonary osteoarthropathy or thyroid acropachy
- Synovial effusion. The joint fluid may be: clear and acellular in association with, for example, serious injury; inflammatory and cellular, as in rheumatoid arthritis; blood-stained, as in traumatic haemarthrosis and bleeding disorders; purulent, as in pyogenic infection or acute crystal synovitis; or milky, as in chylous arthritis associated with filariasis
- Synovial proliferation, as in rheumatoid arthritis
- Rarely, malignant changes, as in sarcoma and secondary carcinoma

Not infrequently, joints are swollen from a combination of two or more of these factors. Swelling of the synovial tendon sheaths or bursae alongside the joints may also contribute greatly to the clinical picture, non-symmetrical involvement of the extensor sheaths on the dorsum of the wrists being very typical of rheumatoid arthritis. Subacromial and semi-membranosus bursal involvement may contribute some swelling when the shoulders or knees are affected by an inflammatory arthritis. If there is doubt regarding the presence of infecting organisms, aspiration should be performed. This will often contribute useful information in non-infective conditions such as gout, chondrocalcinosis or haemarthrosis if the diagnosis is in doubt.

Joints affected by any disease process may show a variable blend of five factors: swelling, pain, stiffness, tenderness and weakness. These five factors in variable combinations cause the dysfunction typical of the particular arthritic disease in question. In progressive systemic sclerosis (scleroderma), stiffness is the dominant component; in gout, swelling, tenderness and pain are predominant. The joints in rheumatoid arthritis vary and differ depending on the activity and stage of the disease process: stiffness in early rheumatoid disease is largely due to joint swelling, and in advanced disease to irreversible change, even, in some patients, to the point of joint fusion. In many instances, chronic joint swelling produces excessive joint laxity, allowing reversible deformity or subluxation. In other cases, the destruction of joint tissue in rheumatoid arthritis causes gross hypermobility: ‘lorgnette’ or ‘telescopic fingers’ are extreme examples of this in psoriatic arthritis.

Two of the cardinal signs of inflammation – heat and redness – are often absent in inflammatory arthritis, while in acute pyarthrosis the joint is hot, and in gout it is hot and red. In most patients with rheumatoid arthritis, the joints tend to be cold and moist without erythema, although swollen, painful and tender. Inflammatory arthropathies usually cause most discomfort in the early morning, the tissues becoming ‘gelled’ with disuse in the night. This early-morning increase in pain and stiffness of the fingers, wrists and shoulders in particular is characteristic of rheumatoid and similar arthropathies; in ankylosing spondylitis, a similar increase in stiffness and pain occurs in the spine, and other affected joints. Painful morning stiffness is also seen characteristically in polymyalgia rheumatica in the shoulder and hip girdles.

There are very many possible causes of joint involvement in systemic disease; these are listed in Box J.3. Not all can be discussed and described, and the following account deals only with the more common and distinctive.

**CONGENITAL ARTHROPATHIES**

In classical achondroplasia, bony growth is abnormal, but epiphyseal development is normal; premature osteoarthritis is not usual, and spinal stenosis may occur. In pseudoachondroplasia and the various spondyloepiphyseal dysplasias, the epiphyses are involved, rendering these subjects particularly prone to severe premature osteoarthritis in adult life. In Conradi’s syndrome, a widespread patchy calcification in articular and other cartilages is seen in infancy, with shortening and asymmetry of the limbs; premature osteoarthritis follows in adult life, as it does in hereditary progressive arthro-ophthalmopathy, in which progressive myopia is associated with multiple epiphyseal dysplasia.
### Box J.3  Arthropathies

#### Congenital
- Achondroplasia and pseudoachondroplasia
- Angiokeratoma corporis diffusum (Fabry's disease)
- Arthrogryposis multiplex congenita
- Camptodactyly
- Chondrodysplasia punctata (Conradi's syndrome)
- Congenital indifference to pain
- Down's syndrome
- Dysplasia epiphysalis multiplex
- Ehlers-Danlos syndrome
- Familial dysautonomia (Riley-Day syndrome)
- Hypermobility syndrome
- Marfan's syndrome
- Morquio–Brailsford syndrome
- Osteodystrophy
- Osteogenesis imperfecta
- Spondylodephysial dysplasia

#### Degenerative, traumatic and occupational
- Ankylosing vertebral hyperostosis (Forestier's disease, diffuse interstitial spinal hyperostosis)
- Occupational syndromes, e.g. porter's neck, wicket-keeper's fingers
- Osteoarthritis
- Traumatic syndromes, e.g. traumatic haemarthrosis

#### Dietetic
- Fluorosis
- Kashin-Beck disease
- Rickets
- Scurvy

#### Endocrine
- Acromegaly
- Diabetic cheiroarthropathy
- Hyperparathyroidism
- Hypoparathyroidism
- Hypothyroid and myxoedematous arthropathy
- Thyroid acropathy

#### Gut-associated
- Acute gastrointestinal bacterial infection
- Antibiotic-induced (pseudomembranous) colitis
- Crohn's disease
- Jejuno-ileal bypass arthritis–dermatitis syndrome
- Ulcerative colitis
- Whipple's disease

#### Idiopathic non-inflammatory
- Acne fulminans arthritis
- Behçet's syndrome
- Dermatomyositis and polymyositis
- Dressier's syndrome
- Erythema multiforme
- Erythema nodosum
- Familial Mediterranean fever
- Henoch-Schönlein syndrome (anaphylactoid purpura)
- Intermittent hydrarthrosis
- Juvenile chronic arthritis (pauciarticular, polyarticular, systemic)

#### Infective
- Mixed connective tissue disease
- Palindromic rheumatism
- Pigmented villonodular synovitis
- Progressive systemic sclerosis (scleroderma)
- Relapsing polychondritis
- Rheumatoid arthritis
- Sarcoïdosis
- Seronegative arthritis, e.g. ankylosing spondylitis, psoriatic arthritis, reactive arthritis, enteroarticular arthritis (Crohn's disease, ulcerative arthritis)
- Sjögren's syndrome
- Systemic lupus erythematosus

#### Idiopathic non-inflammatory
- Osteochondritis dissecans
- Osteochondrosis

#### Haematological
- Agammaglobulinaemia
- Haemophilia and allied disorders
- Leukaemia
- Sickle-cell disease
- Thalassaemia

#### Infections due to bacteria, spirochaetes and Mycoplasma
- Anthrax
- Brucella abortus and melitensis arthritis
- Cat-scratch fever
- Diphtheria
- Erysipelas
- Glanders
- Clutton's joints
- Haverhill fever
- Infective endocarditis
- Jaccoud's arthropathy
- Pyogenic (staphylococcal, psittacosis, gonococcal, pneumococcal, etc.)
- Rat-bite fever
- Rheumatic fever
- Secondary syphilis
- Streptococcal reactive arthritis
- Tuberculosis
- Typhoid and paratyphoid fever
- Weil's disease (Leptospirosis icterohaemorrhagica)
- Pseudomonas pseudomallei (meliodosis)
- Mycoplasma pneumoniae
- Poncet's disease
- Lymphogranuloma venereum
- Meningococcal fever
- Leprosy
- Lyme arthritis
- Yaws

#### Infections due to viruses
- Chikungunya
- Dengue fever
- Echo virus infection
- Glandular fever (infectious mononucleosis)
Arthrogryposis multiplex congenita is a rare congenital condition characterized by joint contractures, usually symmetrical and multiple, affecting the lower limbs rather than the upper, and distal joints more than proximal ones. The subcutaneous tissues may be thick, doughy or gelatinous. While usually apparent at birth and readily recognized, if the patient is not seen until adult life, the condition may be confused with advanced rheumatoid arthritis with joint contractures of the fingers, knees, elbows and wrists. It is usually, but not always, painless, although secondary degenerative changes may occur later and cause considerable discomfort, particularly in the hips, which may become dislocated. Other congenital abnormalities may be present, such as small or absent patellae, a high palate, hypospadias and micrognathia.
Clubbing of the fingers and toes is common, but mental disability is rare.

**Camptodactyly** is an innocent congenital condition where the little fingers are flexed with thickening of the proximal interphalangeal joints. It is a relatively common condition, and important only in that it may be confused with osteoarthritis. Other fingers are less often affected.

In **congenital indifference to pain**, the patient repeatedly traumatizes the joints and other tissues, and may develop secondary traumatic osteoarthritis or even neuropathic joints. Such patients are usually otherwise mentally and neurologically normal, but they are constantly suffering the effects of injury, fractures, bruises, dislocations, cuts and scratches, as they do not experience pain in the same way as normal subjects. It is a rare condition. **Dysplasia epiphysialis multiplex** is inherited as an autosomal dominant trait. The epiphyses of the long bones become deformed, the hips most commonly. Any or all of the epiphyses of the long bones may be affected, with resultant osteoarthritis, particularly of the hips and knees, beginning in early life. The sufferers are often of short stature with short, squat digits. Such children presenting with pain, usually in the hip, are often misdiagnosed as cases of Perthe's disease.

**Ehlers–Danlos syndrome** is a rare genetically determined disorder of connective tissue, characterized by hypermobile joints and hyperextensible skin that tends to split if mildly injured, with resulting gaping scars. There are probably seven entities included in this title; in one, in particular, the ecchymotic type IV, sudden death may occur from spontaneous arterial rupture or gastrointestinal perforation, although abnormalities in collagen biosynthesis occur also in three other types. Joint subluxations and dislocations occur if the joints are hypermobile, and effusions into knees are common. Dislocations of the clavicles, patellae, shoulders, radii and hips may occur and recur several times. Other developmental abnormalities are common, such as kyphoscoliosis, anterior wedging of vertebrae, spina bifida occulta, club foot and genu recurvatum. Bleeding may occur in the superficial tissues, or from the vagina, rectum or mouth, and haemarthrosis may occur. Children may be late to start walking and develop a tabetic-like gait. In the sixth to ninth months of pregnancy, joints may become more lax and subluxable than previously. Premature osteoarthritis may occur in a few patients.

**Familial dysautonomia** (Riley–Day syndrome) is a congenital disorder almost completely confined to Ashkenazi Jews, transmitted as an autosomal recessive trait. Among many manifestations, a relative insensitivity to pain may lead to neuropathic joints in the knee or shoulder in early adolescence.

Maldevelopment of the axial and peripheral joints is a feature of the **mucopolysaccharidoses**. Features of these inherited disorders are due to an accumulation of mucopolysaccharides (or glycosaminoglycans) in the tissues as a result of a deficiency of lysosomal enzymes necessary for their degradation. In the most common of these rare disorders, Hurler's syndrome (gargoylism), the onset is from 6 months to 2 years of age, but the typical picture of gargoylism does not appear until 4 or 5 years later. The child develops coarse features with thick lips, a large bulging head and a flattened nose; the cervical spine is short, with kyphosis of the lower dorsal and upper lumbar regions. The acetabula are shallow and the epiphyses flattened, irregular and retarded in development.

Limitation of joint movement is common, particularly abduction of shoulders and hips and extension of fingers, and contractures of hips and elbows may occur, the hands becoming clawed. The children are often intellectually impaired, and rarely live beyond 20 years of age. Similar features are found in other mucopolysaccharidoses, such as Hunter's syndrome, Sanfilippo's syndrome, Scheie's syndrome and Maroteaux–Lamy's syndrome.

In **type IV Morquio–Brailsford mucopolysaccharidosis**, the children appear normal until 1–2 years of age, when kyphosis is seen, with protrusion of the sternum and prominence of the chin appearing a year later. Growth usually ceases at 10 years, the child showing a short trunk with kyphosis, knock-knees, flat feet, a waddling gait, muscle weakness and increased laxity of the joints, which is in sharp contrast with the contractures seen in the other mucopolysaccharidoses. In a severe case, almost every joint may be affected, the spine, hips and knees most often, and cord compression may occur. Aortic regurgitation is common. Keratan sulphate excretion in the urine is increased and, although not invariably present, may be diagnostic. Mentally, the children are normal.

**Ganglioside storage diseases** also result from inherited enzyme deficiencies. **Farber's disease** results from the deposition of lipid with a vigorous granulomatous reaction (disseminated lipogranulomatosis). Arthritis is an early feature, with red swollen joints and periarticular pigmented swelling appearing in the first few months of life. It is a very rare condition. Death from respiratory infection is usual before the age of 2. In **hypermobility syndrome**, generalized joint laxity occurs as an isolated finding, (Fig. J.20). Symptoms,
more common in females than in males, usually start from the age of 15. The knees are most commonly affected. Degenerative changes may begin early in the fourth decade. These children, who often consider themselves to be ‘double-jointed’, often suffer quite severe cramp-like pains in the legs after sporting activities. About 10 per cent of otherwise normal subjects have such hypermobility. Symptoms are related to the degree of hypermobility but often decrease with advancing age, and crippling osteoarthritis is uncommon.

The picture of Marfan’s syndrome is of a tall, thin loose-jointed youth or girl with long extremities, especially the fingers (arachnodactyly), dislocated lenses, tremulous irides and cardiovascular abnormalities, particularly dilatation of the ascending aorta and aortic regurgitation. Fifty per cent of these patients present with backache, pains in the joints and/or effusions. Joints, most commonly the hips or shoulders, may readily dislocate, but the early development of osteoarthritis is unusual. Many other abnormalities may be present, particularly pigeon chest or pectus excavatum. The sexes are affected equally. The distance from pubis to sole exceeds that of pubis to vertex, and arm span is greater than height. Distal bones are longer than proximal, and subcutaneous fat is sparse.

Osteodysplasty (Melnick–Needles syndrome) is another inherited skeletal dysplasia, probably a congenital disorder of skeletal growth, which leads to early degenerative changes in the large weightbearing joints, including the spine. Radiographs show curvature of the long bones of the limbs, with an irregular cortex and widening and thinning of the metaphyses. These changes may roughly resemble those of rickets, but the ribs, clavicles and scapulae are also deformed. The outstanding abnormality in osteogenesis imperfecta is the ease with which bones may be fractured. In addition, the joints are unduly mobile, thinness of the sclerae gives the eyes the typical pale blue appearance, and there is atrophy of the skin with a tendency to subcutaneous haemorrhage. Joints dislocate readily, and growth may be arrested by multiple small fractures in the epiphyses. Spine and chest deformities may occur, and deafness is not unusual. Ligaments are weakened, and tendon ruptures may occur.

This list of congenital non-infective arthropathies is not complete, and the different causes are not yet clear enough to classify them all accurately. Undue laxity, with recurrent subluxations or increased friability or fragility of the tissues, leads to premature degenerative changes in the joints affected. Other co-existent congenital abnormalities are common.

DEGENERATIVE ARTHROPATHIES

By osteoarthritis is meant the various painful syndromes arising primarily from degenerative changes in the joints (Figs J.21 and J.22). Age brings degenerative changes, but the pains experienced vary greatly depending on psychosocial factors, degree of change and joints affected. Such changes are more likely to occur in any joint previously injured by fracture or dislocation, or even mild subluxation or in any joint that is congenitally abnormal or, because of mechanical factors, working abnormally in the face of abnormal stresses. Repeated ‘microtrauma’ may also predispose to degenerative changes. Endocrine factors may play a part in some cases, as in the so-called generalized osteoarthritis that usually affects women 1–5 years or more before or after the menopause. The most commonly affected joints are the distal and proxima interphalangeal joints of the fingers (Heberden’s and Bouchard’s nodes), the thumb bases (carpometacarpal joints of the thumbs), the cervical and lumbar spine, the knees and hips, and
the acromioclavicular joints. Proximal interphalangeal involvement is common while metacarpophalangeal involvement is uncommon unless secondary. Although essentially degenerative, Heberden’s and Bouchard’s nodes may, in some cases, be tender, red and inflamed in the early stages; inflammation also occurs in osteoarthritis in other joints, but does not dominate the scene as it does in rheumatoid arthritis. The hand in osteoarthritis differs from that in rheumatoid arthritis, as shown in Table J.5.

In the knee, effusions may occur as a result of trauma, and there may also be considerable synovial proliferation. It is not always easy to distinguish an osteoarthritic joint from a rheumatoid one, but the pattern of the disease elsewhere in the body and the absence of systemic features in osteoarthritis usually suffice to distinguish the two. In any joint, the absence of inflammatory swelling, nodules favours osteoarthritis rather than inflammatory arthritis. Sedimentation rates are rarely elevated above 30 mm in the first hour (Westergren) in osteoarthritis and are usually normal, as are haemoglobin and plasma proteins. Tests for rheumatoid factor and anti-CCP (cyclic citrullinated peptides) antibodies are negative. In the cervical spine, degenerative changes centre essentially around the lower fifth, sixth and seventh intervertebral discs, the pain often fanning up into the occiput, over the head and into the shoulders; neck movements are restricted and painful. In rheumatoid disease, particularly in childhood, involvement is more diffuse throughout the cervical spine, although pain may be referred in a similar manner. Subluxation of the first on the second vertebra, which occurs in rheumatoid arthritis and ankylosing spondylitis, is not seen in osteoarthritis. A lateral radiograph of the cervical spine will readily distinguish the rough irregularity of disc degeneration from the straight, even intervertebral bridging of ankylosing spondylitis.

The condition known as ‘ankylosing hyperostosis’ of the spine (Forestier’s disease) is a degenerative one occurring not infrequently in diabetics (see BACK, PAIN IN, p. 45), usually in the lower dorsal area.

**DIETETIC ARTHROPATHIES**
Kashin–Beck disease is a condition apparently resulting from the fusarial infection of flour. In the valleys in eastern Siberia, northern China and North Korea, where it occurred, degenerative changes in the joints and spine appeared in relatively young people as a result of cartilage destruction. It is now disappearing with the elimination of infected grain. Rickets is much less common than it was, and for this reason may be more easily missed. Presenting symptoms may be pains and tenderness over the bones, particularly the back, hips, thighs and legs generally. The pains are usually aggravated by rising from resting positions and by exercise. The dangers of missing the diagnosis lie in permanent bony deformities in the pelvis and lower extremities, and in the thorax. Pelvic deformities, most serious in females, usually occur in the first year of life.

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**Figure J.22** Pelvic radiograph showing osteoarthritis changes in the right hip joint, joint space narrowing, subchondral sclerosis, osteophytes and pseudocyst formation.

**Table J.5 Differential diagnosis of osteoarthritic and rheumatoid joints**

<table>
<thead>
<tr>
<th>Osteoarthritis</th>
<th>Rheumatoid arthritis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bony joint swelling</td>
<td>Spindle soft-tissue swellings of joints</td>
</tr>
<tr>
<td>Terminal interphalangeal joint involvement common (Heberden’s nodes)</td>
<td>Terminal interphalangeal joint less commonly involved</td>
</tr>
<tr>
<td>Metacarpophalangeal and proximal interphalangeal joints less commonly involved</td>
<td>Metacarpophalangeal and proximal interphalangeal joints commonly involved</td>
</tr>
<tr>
<td>Wrists rarely affected</td>
<td>Wrists commonly affected</td>
</tr>
<tr>
<td>Tendon sheaths not involved</td>
<td>Swelling of tendon sheaths common</td>
</tr>
<tr>
<td>Affected joints not usually very tender</td>
<td>Affected joints usually tender</td>
</tr>
<tr>
<td>Gross deformity rare</td>
<td>Joint effusion common</td>
</tr>
<tr>
<td>Radiographs show:</td>
<td>Radiographs show:</td>
</tr>
<tr>
<td>• Juxta-articular bony sclerosis</td>
<td>• Juxta-articular osteoporosis</td>
</tr>
<tr>
<td>• Diffuse joint space loss</td>
<td>• Bony erosions</td>
</tr>
<tr>
<td>• Juxta-articular pseudocysts</td>
<td>• Subluxation or deformity</td>
</tr>
<tr>
<td>• Osteophytes</td>
<td></td>
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</tbody>
</table>
It is associated with hypocalcaemia, cataracts, fits, calcification is present, the sacroiliac joints are normal. Of ankylosing spondylitis but, although ligamentous stiffness may cause a clinical picture similar to that of the fixed bent spine, that of advanced ankylosing spondylitis resembles more that of osteoarthritis or, because of the condition not infrequently occurs, and arthralgia is a common complaint. Signs of inflammation are absent, but synovial stiffness and, occasionally, effusions may occur, the knees and hands being most commonly affected. Traumatic lesions are not uncommon, and hip pain may be due to a slipped femoral epiphysis.

In the original description of myxoedema, in 1873 by Sir William Gull, muscular stiffness, joint swelling and broad spade-like hands were noted. In the early stages of the disease, before the classical features of myxoedema appear, the hands may be mistaken for those of early rheumatoid arthritis. Carpal tunnel syndrome not infrequently occurs, and arthralgia is a common complaint. Signs of inflammation are absent, but synovial stiffness and, occasionally, effusions may occur, the knees and hands being most commonly affected. Traumatic lesions are not uncommon, and hip pain may be due to a slipped femoral epiphysis.

In hyperparathyroidism, as in some cases of osteomalacia, crush lesions may occur in juxta-articular bone with a traumatic type of synovitis, with effusions and impaired function of the affected joints. Calcification is not uncommon in the synovial membrane and cartilage in these cases, but it is rare in rheumatoid arthritis, a point of distinction between the two conditions. Not only may ‘pseudorheumatoid arthritis’ occur, but also ‘pseudogout’ due to deposition of crystals of calcium pyrophosphate dihydrate.

In idiopathic hypoparathyroidism, back pain and stiffness may cause a clinical picture similar to that of ankylosing spondylitis but, although ligamentous calcification is present, the sacroiliac joints are normal. It is associated with hypocalcaemia, cataracts, fits, tetany and rashes. In thyroid acropathy, subperiosteal thickening is seen in the metacarpal and phalangeal bones of the hands in patients with hyperthyroidism who have in many cases been treated and rendered euthyroid. Exophthalmos is common, and so-called ‘pretibial myxoedema’ may be present. The somewhat thickened hand resembles that seen in hypertrophic pulmonary osteoarthritis, but the condition is milder and less extensive, being usually confined to the hands.

ENDOCRINE ARTHROPATHIES

In acromegaly, recurrent pains in the spine and limb joints may be mistaken for those occurring in rheumatoid or osteoarthritis. Effusions may occur in the knees, and carpal tunnel syndrome is not uncommon. The joints may be hypermobile in the early stages due to enlargement of the cartilages, and subluxations and traumatic effusions may occur as in the other hypermobility syndromes. Later bony overgrowth restricts movement, so that the picture resembles more that of osteoarthritis or, because of the fixed bent spine, that of advanced ankylosing spondylitis.

Diabetic cheiroarthropathy affects the hands of some patients with diabetes mellitus; the fingers may become stiff and partially flexed, with waxy thickening of the skin of the palms.

In the original description of myxoedema, in 1873 by Sir William Gull, muscular stiffness, joint swelling and broad spade-like hands were noted. In the early stages of the disease, before the classical features of myxoedema appear, the hands may be mistaken for those of early rheumatoid arthritis. Carpal tunnel syndrome not infrequently occurs, and arthralgia is a common complaint. Signs of inflammation are absent, but synovial stiffness and, occasionally, effusions may occur, the knees and hands being most commonly affected. Traumatic lesions are not uncommon, and hip pain may be due to a slipped femoral epiphysis.

In hyperparathyroidism, as in some cases of osteomalacia, crush lesions may occur in juxta-articular bone with a traumatic type of synovitis, with effusions and impaired function of the affected joints. Calcification is not uncommon in the synovial membrane and cartilage in these cases, but it is rare in rheumatoid arthritis, a point of distinction between the two conditions. Not only may ‘pseudorheumatoid arthritis’ occur, but also ‘pseudogout’ due to deposition of crystals of calcium pyrophosphate dihydrate.

In idiopathic hypoparathyroidism, back pain and stiffness may cause a clinical picture similar to that of ankylosing spondylitis but, although ligamentous calcification is present, the sacroiliac joints are normal. It is associated with hypocalcaemia, cataracts, fits, tetany and rashes. In thyroid acropathy, subperiosteal thickening is seen in the metacarpal and phalangeal bones of the hands in patients with hyperthyroidism who have in many cases been treated and rendered euthyroid. Exophthalmos is common, and so-called ‘pretibial myxoedema’ may be present. The somewhat thickened hand resembles that seen in hypertrophic pulmonary osteoarthritis, but the condition is milder and less extensive, being usually confined to the hands.

GUT-ASSOCIATED ARTHROPATHIES

The more characteristic of these are considered in the next section under the spondyloarthropathies.

IDIOPATHIC INFLAMMATORY ARTHROPATHIES

In dermatomyositis and polymyositis, minimal or moderate transitory arthralgia or arthritis occurs in about one-third of cases. Effusions are less common. The fingers and knees are most commonly affected. The skin and muscle manifestations point to the true diagnosis, muscles of the pelvic girdle, thighs and shoulder girdle becoming weak. The association of dermatomyositis with malignant disease in adult cases should be kept in mind.

Dressler’s syndrome following myocardial infarction or cardiac injury or surgery occurs around 2-4 weeks or more after the acute episode, with pericarditis, arthralgia and, rarely, arthritis.

The diagnosis erythema multiforme probably covers several different entities, some mild, some severe, the so-called Stevens-Johnson syndrome being a severe variant. Arthritis or arthralgia may occur, along with other inflammatory reactions in the skin, eye, mouth and elsewhere.

Familial Mediterranean fever is an ill-understood disorder characterized by recurrent and sometimes periodic attacks of arthralgia or arthritis. It occurs predominantly in people of Mediterranean origin, Armenians, Arabs and Sephardic Jews. The onset is in childhood or adolescence, episodes of fever recurring with polyserositis, abdominal pain, urticaria and other rashes, arthralgia and arthritis and, later, amyloidosis. Joint manifestations occur in one-third to one-half of the cases, usually arthralgia but sometimes mono- or oligoarthritis. The acute episodes last only a few days, rarely weeks, most cases showing no permanent sequelae. Sacroiliac changes may occur late in the disease.

Henoch–Schönlein syndrome (‘anaphylactoid purpura’) is most common in children under 12 years of age. The outstanding feature is a maculopetechial and sometimes papular rash, particularly on the buttocks.
and extensor surfaces of the lower limbs. Urticaria and purpura may occur. Pain, swelling and stiffness of the joints, most commonly the ankles and knees, are usually transient and last only a few days. Alimentary haemorrhage and haematuria are not uncommon, and about 10 per cent of cases develop renal failure.

**Palindromic rheumatism** is a name given to recurring episodes of arthritis due to many causes, the most common probably being the early phase of rheumatoid arthritis.

**Pigmented villonodular synovitis** presents as a persistent but usually relatively painless synovial proliferation with blood-stained joint fluid. Brown nodular masses, possibly due to haemangiomas, form in the synovia; these become traumatized, inflamed and hyperplastic, the hyperplastic synovial cells containing haemosiderin. The condition is usually monarticular, commonly of the knee, and occurs in young adults, males rather than females. The aspirated joint fluid is characteristically blood-stained or dark brown in colour.

In **progressive systemic sclerosis** (scleroderma), the skin is stretched tight over the underlying tissues, the joints being intact, although initially showing changes resembling those of rheumatoid arthritis.

**Relapsing (or atrophic) polychondritis** is a rare disorder in which the cartilages of the joints, ears, nose and trachea soften and collapse; this leads to arthritis, facial changes, dyspnoea or stridor and, occasionally, death.

**Rheumatoid arthritis** (Fig. J.23) is sufficiently well known as to need no description. It is as well to remember that tendons, tendon sheaths and bursae are commonly involved by the inflammatory process, and these add to the clinical picture. **Juvenile arthritis** is, in only a small minority of cases, an early form of rheumatoid arthritis. In a few cases, particularly in boys, it may be an early form of ankylosing spondylitis, but it is usually a seronegative chronic arthritis. It differs from adult polyarthritis in that splenomegaly and lymphadenopathy are more common, tests for rheumatoid factor are usually negative, involvement of the terminal interphalangeal joints of fingers and cervical spine are more common, and skin rashes of the maculopapular type are more common. In the eye, iritis with band opacity in the cornea occurs, sometimes with secondary cataract formation; these are not seen in rheumatoid arthritis in adults. Growth in general may be arrested if the disease is severe, and premature fusion may occur in epiphyses adjacent to the involved joints. Pericarditis is more common in juvenile arthritis than in adult rheumatoid arthritis.

In **Felty’s syndrome**, splenomegaly, enlargement of the lymph nodes, neutropenia and sometimes pigmentation of the skin are superimposed on the usual picture of rheumatoid arthritis. The only reason for maintaining the title in what is merely a variant of rheumatoid arthritis is to emphasize the importance of the neutropenia, for intercurrent infections are the rule, and splenectomy may be necessary. Leg ulcers are relatively common, the usual site being the lower shin anteriorly.

The arthropathy associated with **sarcoidosis** is often accompanied by erythema nodosum; a weak or negative tuberculin reaction is usual, and the arthropathy may be no more than a migratory arthralgia, or it may be a true polyarthritis with pain, fever, systemic upset and swelling of several joints, usually the larger ones and particularly the ankles. In the majority of cases, the polyarthritis subsides in a few weeks. Hilar node enlargement is common in chest radiographs, and lymph nodes may be palpable in the neck and axilla in some cases. Splenomegaly may be present. Histoplasmosis may also present with hilar lymphadenopathy and joint pains (‘pseudosarcoidosis’).
Joints, Afections of

The term *spondyloarthritis* is applied to a family of conditions whose key features are: involvement of the spine and sacroiliac joints; oligoarticular asymmetrical arthritis; an association with iritis, psoriasis and inflammatory bowel disease; and a high prevalence of the HLA B27 antigen. The principal members of this group are ankylosing spondylitis, reactive arthritis, psoriatic arthropathy, and enteropathic arthritis associated with ulcerative colitis and Crohn’s disease.

The classical picture of *ankylosing spondylitis* is that of a young male adult with a stiff back, sometimes with thoracic or cervical spine or hip involvement. The erythrocyte sedimentation rate may be elevated, anterior uveitis is present in 25 per cent of cases at some stage in the disease course. Changes on plain radiographs are often a late sign and if there is clinical suspicions MRI should be used. Peripheral joint involvement may occur. Knee effusions are not uncommon. The pattern of the disorder is essentially the central, spine and girdle joints being predominantly affected, and the peripheral small joints less frequently; this contrasts with rheumatoid arthritis. Nodules do not occur, and rheumatoid factor is not present in the serum. The histocompatibility antigen HLA-27 is found in over 90 per cent of patients.

Reactive arthritis comprises arthritis associated with genital tract inflammation or recent gastrointestinal infection. The syndrome may be caused by either sexually transmitted infection or acute gastrointestinal infection and, in either case, urethritis or cervicitis may be present. Recognized causal pathogens include *Chlamydia trachomatis*, *Salmonella enterica* Enteritidis and Typhimurium, *Shigella flexneri*, *Yersinia enterocolitica* and *pseudotuberculosis*, and *Campylobacter jejuni*. A reactive arthritis may also be seen after streptococcal throat infection. Traditionally, arthritis, urethritis and conjunctivitis comprise the classical triad but this combination is rare; conjunctivitis is often transient or mild, and genital tract symptoms may be mild, overlooked or denied. Diagnosis therefore requires a careful history and a genitourinary examination, including a microscopic examination of urethral and/or cervical smears.

Arthritic symptoms appear a few days or up to 3 weeks after the initial symptoms of the causative infection. The distribution of affected joints, ankles, heels and knees being principally affected, is characteristic, and lesions of the buccal mucosa, the glans penis or prepuce (balanitis circinata), or skin (keratoderma blennorrhagica, although this is very rare) suggest the correct diagnosis. Later, sacroiliac changes may occur, and sometimes a clinical picture similar to that seen in ankylosing spondylitis develops. Rheumatoid factor is absent from the blood; nodules do not occur. When skin manifestations are present, the condition may closely resemble that of psoriatic arthropathy. The interphalangeal joints of the toes, less frequently involved in rheumatoid arthritis, may be affected in reactive arthritis. Iridocyclitis and iritis, rare in rheumatoid arthritis, are not uncommon.

In *psoriatic arthropathy* (Fig. J.24), the arthritis usually but not invariably follows the skin disorder by several years: the diagnosis can be made in the absence of skin disease particularly if there is a history in a first-degree relative. The seronegative polyarthritis tends to be more patchy and less evenly symmetrical than that of rheumatoid arthritis, and the terminal interphalangeal joints of the fingers are frequently affected, particularly if the nails are affected by the pitting, ridging and separation of psoriasis. Dactylitis presents as diffuse inflammation of a digit due to joints and tendon inflammation. In some cases, the sacroiliac joints or the spine are affected, the clinical picture being that of ankylosing spondylitis.

In *Crohn’s disease, ulcerative colitis* and, less commonly, *Whipple’s disease* (intestinal lipodystrophy), arthralgia or arthritis may occur in the spine or peripheral joints. In all of these, rheumatoid factor is absent from the blood, and rheumatoid nodules are not seen. In the arthropathy of ulcerative colitis, the best documented of these three disorders, the onset is usually between the ages of 15 and 45 years. It is usually symmetrical and often monarticular, with short exacerbations and usually complete recovery, joint erosions being rare and minor in character. It affects both sexes equally,
and usually begins acutely, affecting one knee or ankle primarily, subsequent attacks being of similar pattern. The arthritis usually commences long after the onset of the colitis, and may coincide with an exacerbation of the disease. In all three conditions—ulcerative colitis, Crohn’s disease and Whipple’s disease—a picture similar to that of ankylosing spondylitis may eventually appear after some years.

In systemic lupus erythematosus (SLE), any or all systems of the body may be involved in addition to the joints, which are not invariably involved, although arthralgia is usually present at some stage in the course of the disease. The patient, usually a female, is more ill than arthritic in most cases, although joint involvement is present in about two-thirds of patients. The finding of numerous antibodies, including antinuclear antibody, in high titre, and dsDNA antibody in the blood is strong confirmatory diagnostic evidence. The joint involvement may be flitting, resembling rheumatic fever, or more constant, resembling rheumatoid arthritis. A deforming but non-destructive arthropathy (Jaccoud’s arthritis) is less common. The co-existence of skin lesions and visceral manifestations suggests the correct diagnosis, the typical lupus butterfly rash over nose and cheeks being particularly characteristic. Neutropenia and anaemia are common, and thrombocytopenia not uncommon. Asthma, proteinuria, neurological signs, splenomegaly, retinal exudates and a number of other co-existent findings in any patient with arthritis should make one think of this disorder or a related connective tissue disease. Epileptiform fits occur in about 10 per cent of cases. Patients with neurological and renal involvement fare worst. Patients having a combination of clinical features of systemic lupus erythematosus, progressive systemic sclerosis and polymyositis with high titres of a circulating antinuclear antibody with specificity for a nuclear ribonucleoprotein are said to have mixed connective tissue disease.

IDIOPATHIC NON-INFLAMMATORY ARTHROPATHIES

In osteochondritis dissecans, flakes of articular cartilage, sometimes with a portion of the underlying bone, become detached without evident trauma, the condition manifesting itself as recurring attacks of arthritis. The most common site (85 per cent) is the knee; the radial head is the next most common, the hip and ankles being rarely involved. The condition may be bilateral, and X-rays are usually diagnostic.

In osteochondrosis, the diagnosis is also essentially a radiological one. It is essentially a disturbance of epiphyseal ossification seen in childhood and early adult life, possibly ischaemic in origin. Early radiographs show dense fragments in the epiphysis and a broadening of the epiphysial line with, later, areas of rarefaction and condensation, so that a core of dense bone is seen in a porotic matrix. The epiphyses are affected during the periods of their greatest activity, for instance the femoral head from 4 to 12 years (Legg–Calvé–Perthes disease), and the tibial tubercle from 10 to 16 years (Osgood–Schlatter disease).

HAEMATOLOGICAL ARTHROPATHIES

It is wise to perform a full blood count, erythrocyte sedimentation rate and examination of plasma proteins in obscure cases of arthritis. Approximately 25 per cent of patients with agammaglobulinaemia, congenital or acquired, develop a non-suppurative arthritis not unlike rheumatoid arthritis, the joints showing effusions, pain, tenderness and stiffness. The condition is usually asymmetrical, is unaccompanied by radiological changes, and may be transient, subsiding in a few weeks without sequela, or may persist for years but with little residual change. Biopsy of synovial tissue does not distinguish between the two conditions. The sedimentation rate is usually normal, and tests for rheumatoid factor are negative. In some cases, arthritis has been attributed to Mycoplasma infection, but recurrent infection with the usual pyogenic organisms is also common.

Haemarthrosis may occur in haemophilia (factor VIII deficiency) and allied disorders, such as Christmas disease (factor IX deficiency), and in patients on anticoagulant therapy, but is rare in von Willebrand’s disease. In leukaemia, haemorrhages are common, and flitting pains resembling rheumatic fever are not uncommon in acute leukaemia; this, taken in conjunction with a systolic cardiac murmur, may cause diagnostic confusion, particularly in acute aleukaemic leukaemia. Pains in the bones and joints occur not infrequently in acute leukaemia in childhood and in chronic leukaemia in adults, both myeloid and lymphatic. In children, juvenile arthritis is often diagnosed in error.

In sickle-cell anaemia, painful crises occur that are characteristic of the disorder, and these may occur not only in the abdomen but also in the bones and joints in children or adults. Although the most common symptoms are those of anaemia, some patients have no complaints except during crises. Aseptic necrosis of bone may occur, particularly in the head of the humerus or femur, radiographs showing subsequent areas of increased density and areas of necrosis.
The course of the disease is that of a chronic haemolytic process punctuated by periodic painful crises. Chronic ulceration of the lower legs is relatively common, and scars are commonly to be seen around the malleoli. Another striking complication of sickle-cell disease, particularly in children, is Salmonella osteomyelitis, often multifocal. In thalassaemia major, pains and swelling may occur in the ankles and feet.

**INFECTIVE ARTHROPATHIES**

In the infective arthropathies, the infecting organism is present in the locomotor tissues; in gonococcal monoarthritis, for instance, gonococci can be isolated from the infected joints or joint; the condition responds to appropriate antibiotics. Any of the infections due to bacteria, spirochaetes or *Mycoplasma* may, if there is destruction of tissue, lead to chronic changes in the bones and joints, but, if the correct treatment is given early, there may be little or no residual disability. In these days of extensive and rapid worldwide travel, conditions previously unknown in residents of one country can occur, with resulting arthralgia or arthritis.

Viral arthropathies are common throughout the world, but these are usually mild and transient. Arbovirus infections including Chikungunya and O’nyong-nyong are common in some parts of Africa and South America. To a lesser extent, arbovirus infections also occur in Scandinavia (Ockelbo and Pogosta) and Australia (Ross River virus arthritis). In Europe and the USA, parvovirus arthritis is the most common viral joint disease, also associated with a transient rash, upper respiratory infection and malaise (erythema infectiosum – fifth disease). Arthritis following natural rubella is uncommon because of widespread immunization against the virus, but it may follow vaccination itself. If viral arthritis is suspected, hepatitis B infection must also be excluded. Joint involvement is usually polyarticular and symmetrical, and carpal tunnel syndrome may develop. Symptoms generally subside within 3 weeks. Post-vaccination arthritis may affect a single joint only and persist or recur. Lyme arthritis, named after the part of East Connecticut in which it was first identified, comprises a variable multisystem disease combined with transient asymmetrical oligoarthritis. The causative agent is a spirochaete, *Borrelia burgdorferi*, which is transmitted by tick bites. The disease is only acquired, therefore, in areas where ticks of the genus *Ixodes* are endemic. The disease responds to antibiotic treatment.

Rheumatic fever is seen much less often today than previously. It is as well to remember that many other arthropathies may present in similar form, joints being successively affected and remitting rapidly, the so-called ‘flitting pains’ rippling round the locomotor system. Not only may rheumatoid arthritis present in this way, but so too can systemic lupus erythematosus, ankylosing spondylitis, Hodgkin’s lymphoma, leukaemia, brucellosis and a number of other disorders. The heart is rarely seriously involved if rheumatic fever first occurs over the age of 17 years. Jaccoud’s arthritis is an extremely rare disorder following repeated attacks of rheumatic fever, characterized by ulnar deviation of the fingers and hyperextension of the proximal interphalangeal joints without bone destruction. It is more frequently seen as a manifestation of systemic lupus erythematosus.

**METABOLIC ARTHROPATHIES**

The most common of these is gout (Fig. J.25). This disorder is characterized by the sudden agonizing nature of the acute attack, which is often so severe as to make the patient, almost always an adult male, feel he must have broken a bone in his foot but for the fact that the disorder frequently starts in bed in the early morning. There are usually clear signs of inflammation, the skin being tense, shiny, hot and red over the big toe metatarsophalangeal joint, ankle or hand, the last named being the most common. Acute attacks may also occur in other joints, particularly the knee and wrist. Although hyperuricaemia is usually present, it is not invariably so, and an elevated plasma urate concentration occurs in many other disorders, and is not in itself diagnostic. The presence of tophi in the ears or elsewhere suggests the diagnosis, although the symptoms and signs are usually diagnostic. The only absolute proof is the identification of urate crystals from the affected joint under the polarizing microscope.

In some cases, suggestive of gout, intra-articular crystals turn out to be not urate but calcium pyrophosphate, the condition being *chondrocalcinosis articularis* or ‘pseudo-gout’. This condition most commonly affects the knees, but other joints are also affected, infrequently in symmetrical fashion. The appearance of calcification in the joint cartilages on X-ray (chondrocalcinosis) suggests the diagnosis. Acute inflammatory episodes occur also in chronic renal failure, with the deposition of calcium salts in the soft tissues alongside, rather than in, the joints.

In *calcinosi circumscripta*, calcium salts (carbonate and phosphate) may be deposited under the skin, but they are again para-articular rather than in the joint tissues, which appear normal.

Although rare these days, *amyloidosis* may be secondary to rheumatoid arthritis, ankylosing...
spondylitis and (more rarely) reactive arthritis, but it may also occur in primary form associated with pains and swellings in the joints and, when associated with multiple myeloma, may cause carpal tunnel syndrome. It may also be associated with periodic fevers such as familial Mediterranean fever.

Joint symptoms and backache in particular occur in **ochronosis**. Here, the diagnosis is made by examination of the urine for homogentisic acid and the cartilage of the ears for pigmentation. Radiographs of the spine are typical, heavy calcification occurring in the intervertebral cartilages.

**Multicentric reticulohistiocytosis (lipoid dermatopathy)** may be mistaken for rheumatoid arthritis in adults as changes in the fingers and tenosynovitis occur, but the presence of yellow nodules on the ears, forehead, neck, forearms and elsewhere, with groups of purple papules, suggests the true diagnosis, which can be confirmed by biopsy. In advanced cases, erosion of the phalanges leads to shortening of the fingers.

In **Wilson's disease**, characterized by the accumulation of copper in the tissues, arthritic changes, most commonly in the hands, wrists and knees, may start around the age of 30. Associated features are hepatic cirrhosis and psychiatric disease, the green-brown Kayser-Fleischer ring around the cornea being diagnostic.

**VASCULAR ARTHROPATHIES**

Avascular necrosis occurs in **Caisson's disease** (nitrogen or air embolism), from fat embolism, and occasionally in chronic alcoholism. The hips are often bilaterally involved, with destruction of parts of the heads of the femurs, but the shoulders and one or both knees may also be affected.

**Giant-cell arteritis** and **polymyalgia rheumatica** are probably two facets of the same condition occurring in the elderly as, on existing evidence, both conditions are due to an arteritis of those vessels having an internal elastic lamina. Renal and cerebral vessels are therefore usually spared. The patients, usually over the age of 60, are of either sex, have marked morning stiffness, erythrocyte sedimentation rates up to 100 mm in the first hour (Westergren), and pains and stiffness of the shoulder and hip girdles. When the temporal vessels are involved, a splitting headache is often present, and the main danger is to vision if branches of the ophthalmic artery become affected. The pulses may disappear, and murmurs may be heard at the points of arterial narrowing. The disorder, as far as the girdle joints in general are concerned, is one of pain and stiffness in the hip and shoulder girdles without progressive clinical or radiological change, and eventually with full recovery. Diagnosis of GCA can be confirmed by arterial biopsy or, if indicated, PET imaging.

**NEOPLASTIC ARTHROPATHIES**

Metastatic malignant disease and **multiple myeloma** usually cause bony rather than joint changes. The serum alkaline phosphatase, often elevated in the former, is usually normal in the latter, as there is no osteoblastic activity in myelomatosis. Joint changes occur, however, in **hypertrophic pulmonary osteoarthropathy**, a condition...
mostly associated with a bronchial carcinoma, usually a peripheral one. Removal of the primary lesion leads to rapid resolution of the effusions and arthritic changes in the more commonly affected joints, the knees and ankles. The fingers and toes are clubbed, and the extremities show a thickening based on new subperiosteal bone deposition, which can be seen in radiographs. Not all cases of hypertrophic osteoarthropathy are secondary to malignancy, however, some being due to cyanotic congenital heart disease, colonic and other conditions (see FINGERS, CLUBBED, p. 200). In a familial primary form, symptoms of pachydermoperiostosis start usually in adolescence, more commonly in males, the hands and feet enlarging, with marked clubbing and cylindrical thickening of the forearms and legs; recurrent joint effusions may occur. The patient’s features thicken, giving a leonine appearance.

Osteoid osteoma is a benign disorder and, although not a disease of joints, it should be mentioned because of the pain it causes and the difficulties in differential diagnosis. The pain is initially intermittent but becomes more persistent and severe, and it is often aggravated by movement. There are no physical signs. It affects adolescents and young adults and, although any bone except the skull may be affected, the most common to be involved are the femur and tibia, which account for half the cases. Radiographs show a characteristic central opacity surrounded by a translucent zone, surrounded in turn by a zone of sclerosis. It may affect the bones of the spine, where it is often very difficult to diagnose and is usually not suspected. The pains are sometimes worse at night than during the day.

NEUROPATHIC ARTHROPATHIES
Although carpal tunnel syndrome is not strictly a joint affection, it is so often a manifestation of rheumatoid arthritis that it should be mentioned. It is infrequently the first sign of this disorder. Other causes are pregnancy, acromegaly, myxoedema, multiple myeloma and amyloidosis, although it is most commonly idiopathic with no apparent cause. Characteristic symptoms are tingling in the fingers in the median nerve distribution less commonly radiating into the arms, interfering with sleep.

Neuropathic joints (Fig. J.26) in the form of Charcot’s joints in tabes dorsalis are characterized by their gross deformity, painlessness and florid X-ray appearances, in which a number of bone islands surround a grossly deformed or disorganized joint. Syringomyelia affects chiefly the shoulder and elbow; the knee, ankle, hip and spine are more commonly affected in tabs. Diabetic arthropathy is different in that clinical and radiological signs of infection are often present, along with poor vascularization and signs of peripheral neuropathy; the condition is usually confined to the feet and toes. Osborne’s syndrome is due to ulnar nerve compression beneath the arcuate ligament just below the elbow.

In the shoulder-hand syndrome, a reflex dystrophy, trophic changes follow soon after injury to the shoulder or weeks or months after myocardial infarction; a similar syndrome has been reported in patients on antituberculous therapy and in other pathological conditions. The shoulder is stiff and painful, the skin of the hand shiny and smooth and sometimes hyperaesthetic, and the muscles atrophic. There is no joint swelling, although initially there may be considerable swelling of the whole hand and fingers. X-rays initially show osteoporosis of the humeral head.
and wrist, and later a more diffuse ‘ground-glass’ appearance. In many cases, there is no apparent cause for the condition.

**DRUG-INDUCED ARTHROPATHY**

Alcoholics are especially likely to sustain injuries to the bones and joints; they are also more prone to septic arthritis and avascular necrosis of bone. Prolonged **corticosteroid therapy** may also be associated with septic arthritis, osteoporosis and fractures. Crush fractures of the lumbar or dorsal vertebrae are not uncommon. Drug-induced systemic lupus erythematosus can be due to a large range of drugs, the most common being procaine amide; this is the so-called *hydralazine syndrome*. Symptoms disappear on stopping the drug. It has also been reported with oral contraceptives, although such cases are very rare.

**MISCELLANEOUS ARTHROPATHIES**

The knuckle pads (Garrod’s pads), seen not infrequently on the dorsal aspects of the proximal interphalangeal joints of the fingers, are usually not accompanied by any symptoms, and are best disregarded. They are due to fibrous thickenings the size of small orange pips, and are not part of the clinical picture of osteoarthritis or any other form of arthritis. They are not associated with any bony changes, although in some cases they occur with *Dupuytren’s contracture*, which, in turn, is occasionally associated with Peyronie’s disease (induration penis plastica). The palmar contractures occurring in rheumatoid arthritis may, on occasion, resemble Dupuytren’s contracture. *Thorn synovitis* is an inflammatory condition due to a thorn or splinter of wood or a foreign body being knelt on by a child, who is hardly aware of it at the time. The *septic focus syndrome* is a rare disorder in which diffuse aches and pains in and around joints are rapidly relieved by the removal of a septic focus or drainage of an abscess. No residual changes are left in the tissues. Lastly, the *xiphoid syndrome* refers to pains that stem from a displaced or mobile xiphisternum, often the result of trauma. This simple condition is only noteworthy in that it may be mistaken for more serious disorders of the stomach, duodenum, gallbladder or heart.

**JOINT DISEASE IN YOUNG CHILDREN**

The following conditions should be considered when children under 5 years of age present with joint symptoms:

- **Septic arthritis**: due to staphylococci, haemolytic streptococci, *Haemophilus influenzae*, tuberculosis
- **Associated with or following infection**: adenovirus, rubella, mumps, chickenpox, *Mycoplasma pneumoniae*, cytomegalovirus, rickettsia, Lyme arthritis, Kawasaki’s syndrome
- **Idiopathic**: chronic juvenile arthritis (Still’s disease), familial Mediterranean fever
- **Vascular and haematological**: Henoch–Schönlein syndrome, sickle-cell disease, leukaemia, haemophilia, haemangioma, hypogammaglobulinaemia
- **Dietetic**: rickets
- **Miscellaneous**: Farber’s disease, the mucopolysaccharidoses (e.g. Hurler–Scheie syndrome), injuries (battered child syndrome), neuroblastoma, thorn synovitis.

**Infecative (septic conditions)**

*Infection in infancy*

Staphylococcal infection is common, but many organisms may be responsible. The infant is ill, often rejecting food, vomiting or convulsing, but sometimes only mildly ill with slight fever. The hip is the most common joint to be affected, and it is held flexed and adducted, oedema appearing around the adductors.

**1–5 years of age**

*Haemophilus influenzae* infection is common in Great Britain. If several joints are affected, suspect hypogammaglobulinaemia or some other immune abnormality. Staphylococcal, haemolytic streptococcal and, more rarely, tuberculous infection should also be considered.

**Other infections**

Other infections with adenoviruses often start with pharyngitis followed a few days later by fever, macular erythematous rash and a symmetrical arthritis that lasts up to 6 weeks. A similar transient arthropathy may occur with rubella, mumps and chickenpox. Infection with *cytomegalovirus* is often associated with abnormal tests of liver function, and infection with *Mycoplasma pneumoniae* with erythema multiforme. Other infections not seen in the UK unless imported are rickettsial infections such as Rocky Mountain spotted fever or Lyme arthritis, in which a small red macule or papule enlarges to form a large erythematous ring followed by fever and arthritis, usually of only a few joints. In Japan and the East, Kawasaki’s syndrome should be considered a possibility.

**Non-infective inflammatory conditions**

Juvenile chronic arthritis often presents under 5 years of age (see above). If of inflammatory onset, it has to be distinguished from the infective conditions described above.
KELOID

Barry Monk

A keloid is a benign but uncontrolled fibrous overgrowth of the dermis in response to injury. The tendency to form keloids is a personal trait, more common in black individuals, in young adults and in the stretched skin of neck and chest (Fig. K.1). Usually, the antecedent damage is obvious, for example surgical incisions, pierced ear lobes (Fig. K.2), burns and chickenpox. Keloids may follow acne and folliculitis, chiefly on the chest (Fig. K.3) and at the back of the neck, where ingrowing hairs may be a perpetuating factor. Apparently spontaneous keloid formation may occasionally occur. Treatment is invariably unsatisfactory and keloids commonly recur following excision, even if intralesional steroids are injected at the time of operation. Care must be taken to avoid unnecessary trauma (e.g. excision of benign moles) in susceptible subjects.

KIDNEY, PALPABLE

Ben Challacombe

Kidneys are generally difficult to palpate. Palpation of the kidney is easier in a thin individual, but is difficult in those who are obese, where even a considerably enlarged kidney may not be palpable. The causes of an enlarged kidney that lead it to be felt on clinical examination are:

- Renal tumours (renal cell carcinoma)
- Large obstructed kidneys (e.g. pelvi-ureteric junction obstruction or calculi)
- Polycystic disease of the kidney (adult polycystic kidney disease)
- Solitary or multiple large renal cysts (Fig. K.4)

It is usually only large swellings of the kidney that are noticed by the patient, because of either an increase in girth (tight-fitting clothing) or abdominal discomfort. In bilateral polycystic kidneys, the abdomen will be symmetrically enlarged. In unilateral

Figure K.1 Keloid following a knife wound.

Figure K.2 Keloid following ear-piercing.

Figure K.3 Spontaneous keloid.
renal enlargement, one side of the abdomen may be
distorted. For most palpable kidneys, even if they are
asymmetrically enlarged, distinction of the features is
difficult. The kidney should initially be palpated from
the front, and then an attempt made to ‘ballot’ the
kidney using one hand posteriorly, behind the lower
ribs, pushing the kidney forwards so that the kidney
can be felt with the anterior examining hand. Rough
palpation of suspected renal tumours should be
avoided to reduce the theoretical risk of dissemination
of tumour cells. Ultrasound or computed tomography
(CT) scanning usually confirms the enlargement and
suggests the diagnosis.

The rapid onset of a varicocele in adult life (in a
matter of weeks), particularly if on the left, should raise
concern over the presence of a renal cancer. Renal
tumours may obstruct the testicular vein where it drains
into the renal vein on the left and inferior vena cava on
the right, giving rise to an acute varicocele (Fig. K.5).
Currently, renal tumours present most commonly as an
incidental finding during abdominal imaging for other
purposes. Other common signs include macroscopic
haematuria, loin pain, an abdominal mass or with
metastatic disease (fracture or bone pain). Suspicion
of a diagnosis of renal tumour will lead to ultrasound
or contrast CT scanning of the urinary tract, when the
diagnosis will be made.

A pelvic kidney, which is found at or below the brim
of the pelvis, may be felt in the iliac fossa. A renal
transplant will also be easily palpable in the iliac fossa
(often on the right). Anomalies of fusion of the kidney,
such as a horseshoe kidney or crossed, fused renal
ectopia, can sometimes be palpated. A suprarenal
tumour may be sufficiently large to be palpable in its
own right, or it may push the kidney down and make it
palpable, or push the liver forwards such that the liver
becomes palpable.

A renal swelling may be so slight that it is only
found upon radiological examination, or it may
be large enough to attract the patient’s attention.
Hydronephrosis, pyonephrosis, renal tuberculosis,
renal abscess and cysts (single or multiple) in the
kidney have to be diagnosed not only from one
another but also from other tumours simulating a renal
swelling.

The classic characteristics of a renal tumour on clinical
examination are listed below:

- The intestine is in front of the tumour: when either
kidney is merely slightly enlarged, both the large
and small intestines will be in front of it, but when
the organ is so enlarged as to reach the anterior
abdominal wall, the coils of small intestine are
pushed aside. The anatomical relationship of the
large intestine to the kidney, and the absence of a
mesentery, reduces the mobility of the colon, which
usually retains its position in front of the kidney.
Hence an area of resonance can usually be obtained
in front of a renal swelling; bowel is almost never
placed in front of a splenic or ovarian tumour.

- The area of dullness to percussion is continuous
from the lateral aspect of the swelling to the midline
posteriorly; i.e. there is no area of resonance
between the mass and the vertebral spines, as with
a splenic or ovarian tumour.

- A renal tumour usually retains the shape of the
kidney: it is rounded at its borders and poles, and
does not possess any edge or sharp margin, as do

Figure K.4 Renal cysts.

Figure K.5 Left renal tumour causing varicocele.
splenic or hepatic swellings (Fig. K.6). The surface of the tumour may present rounded, smooth, raised bosses in cases of renal growths, or in polycystic disease.

- A renal tumour in the process of enlargement projects forwards and downwards. It may fill up the natural hollow of the loin, but seldom causes any prominence posteriorly. A perinephric abscess, which often simulates a renal swelling, may cause a distinct prominence in the loin.

- A renal tumour may be movable downwards or inwards, unless it is fixed in the loin by preceding inflammation or by the spread of carcinoma into the peri-renal tissues; an enlarged kidney may be felt bimanually and, if grasped between the two hands, can be pushed into the loin. A renal tumour rarely descends into the iliac fossa, but it may be present there in congenital ectopia or in cases of excessive mobility.

- When a renal tumour is large enough to reach the anterior abdominal wall, it commonly comes into contact with it at the level of the umbilicus, at the same time bulging the iliacostal space outwards. There is usually a line of resonance between the upper margin of the tumour and the hepatic dullness.

- A varicocele may be developed on the same side as the renal tumour due to obstruction of the testicular vein, which drains into the renal vein on the left or directly into the inferior vena cava on the right. This is especially significant on the right side, although it is a rare finding.

- With a renal tumour, there may be changes in the urine pointing to renal disease, intermittent painless haematuria (see HAEMATURIA, p. 241), but the urine at any one time may be normal and free from blood or pus. This may be due to the fact that the ureter on the diseased side is blocked, or that the disease does not involve the renal pelvis.

- In exceptional cases, a tumour of the right kidney may extend upwards towards the dome of the diaphragm, rotating the liver so that the anterior margin of the latter descends below the costal margin and prevents satisfactory palpation of the renal areas.

Although, from the above physical characters, it would seem that a renal tumour should present little difficulty in diagnosis, it is by no means infrequent to find that a tumour possessing several of these characters may give rise to considerable doubt in the determination of the organ from which it arises. The following points will assist in the diagnosis of renal swellings from other tumours with which they are likely to be confused.
THE DIFFERENTIAL DIAGNOSIS OF ABDOMINAL
AND PELVIC ORGAN ENLARGEMENTS

Enlargement of the gallbladder
This is placed immediately below the costal margin, so that no interval exists between the tumour and the lower margin of the liver. It is usually oval in outline, with the long axis in the line between the ninth right costal cartilage and the umbilicus, and it is freely movable with respiration and movable from side to side about an axis at the costal margin. There is dullness on percussion over it, and it cannot be felt in the loin or grasped bimanually. With an enlarged gallbladder, there may be attacks of colic, with or without jaundice. Ultrasonography is a particularly valuable non-invasive method of demonstrating the distended gallbladder, and it will also show up gallstones, which are highly echogenic.

Enlargements of the liver
These pass downwards from beneath the costal margin so that there is no line or resonance, or area in which the hand can be depressed, between the tumour and the costal margin. Hepatic tumours do not impair the normal resonance in the loin in the same manner as a renal tumour does. A tongue-shaped lobe of the liver (Riedel's lobe) may cause difficulty in diagnosis, but here the lower margin is seldom as rounded as that of a renal tumour, nor will the mass be felt in the loin on bimanual examination. A tumour or cyst in the concave aspect, or of the left lobe, of the liver is especially liable to cause error in diagnosis, whereas a tumour of the right kidney that projects upwards behind the liver may so rotate the latter that its anterior margin descends below the costal margin and completely obscures the kidney. In a case of a large carcinoma of the right kidney, the liver may in this way be so depressed as to render palpation of the kidney impossible. Ultrasonography (and CT scanning) will readily differentiate between a hepatic and a renal swelling, but an intravenous urogram (IVU) examination may reveal a normal renal picture or a hydrenephrosis or renal mass, and it is generally now not performed in this situation.

Enlargements of the spleen
These descend from beneath the left costal margin and have no bowel in front of them; they are therefore dull to percussion. The edge of a splenic tumour is usually well defined and often notched, and there is resonance between the posterior aspect of the tumour and the spinal column. A splenic tumour is more movable than a left renal tumour. A blood count may help in deciding in favour of a splenic enlargement, and ultrasonography or CT scanning readily differentiates between the two organs.

Perinephric effusions
Blood, pus or urine may form a mass in the loin that may, on examination, be mistaken for a renal swelling. A perinephric effusion may arise from suppuration of the kidney, so that the previous history and examination of the urine will not assist in the differential diagnosis, or it may be due to conditions entirely distinct from renal disease. An effusion of blood around the kidney is, in nearly all cases, caused by trauma to the loin and will be accompanied by other signs of injury. It may, however, occur from the spontaneous growth and rupture of a renal neoplasm or cystic renal disease. A perinephric abscess forms a less well-defined mass than those caused by true renal swellings, and it is associated with signs of sepsis, such as pain, tachycardia and pyrexia. The skin over it may be thickened or oedematous, and fluctuation may be felt to be more superficial than in a renal swelling. A perinephric abscess may result from a carcinoma or diverticulum of the large bowel, from appendiceal inflammation, or from suppuration in a perinephric haematoma due to injury; it may be a sequel to a specific blow, or be due to a haematogenous infection.

Tumours arising from the pelvic organs
Tumours arising from the pelvic organs, from the ovary or uterus, may in some cases simulate renal tumours. An ovarian cyst with a long pedicle occupying the loin may be mistaken for an enlarged or movable kidney, and any sudden attacks of pain occurring from torsion of the pedicle may be confused with renal colic. The normally placed ovarian cyst or uterine fibroid will seldom be confused with a renal swelling, as it is placed in the midline of the body, can be felt to come up from the pelvis, and can be felt on bimanual vaginal examination to be attached to the uterus or its appendages. These tumours give rise to dullness anteriorly and do not alter the normal resonance in the loin. In cases of malignant ovarian tumours associated with ascites, the lumbar resonance may be lost, but on turning the patient over on one side, the previously dull note becomes replaced by resonance in the uppermost loin. In the case of an ovarian cyst with a long pedicle, or of a uterine fibroid of pedunculated, subserous form, the position in the loin may sometimes suggest a renal tumour; it will be found, however, to occupy a more anterior position in the abdomen than a renal tumour and to possess a much greater range of movement, and it does not slip back into the loin under the costal margin in the same manner as an enlarged
kidney does. There is also resonance posteriorly, the kidney as well as the abdominal tumour may be actually palpated, while a distinct connection with the pelvic organs can sometimes be traced from the tumour when the latter is drawn up.

In contrast to the above, a very large cystic renal swelling may be mistaken for an ovarian cyst. It may occupy the greater part of the abdomen, and even be felt per vaginam to be encroaching upon the pelvis; however, on careful examination of a renal tumour of this form, there will be no line of resonance between the mass and the vertebral column posteriorly, the natural hollow of the loin will be filled up, and there is frequently a distinct bulging in the lower thoracic wall, together with an increased length of the iliacostal space on the affected side. Ultrasonography and/or CT scanning will usually enable an accurate anatomical diagnosis of the pelvic mass.

Suprarenal tumours
Suprarenal/adrenal tumours may occasionally be of sufficient size to form an abdominal tumour, presenting as a rounded, movable swelling in the hypochondrium.

Faecal accumulation in the colon, caecum or sigmoid flexure
This may give rise to a tumour and pain of a colicky nature in the loin; the examining fingers can sometimes indent the tumour. They will be distinguished from renal swellings by the general intestinal symptoms, flatulence and the changes in form consequent on the administration of large enemas. A patient with a collection of faeces in the colon may not complain of constipation but may in fact have a small daily evacuation from the overloaded bowel (overflow diarrhoea).

Appendicular inflammatory mass
This will be diagnosed from renal tumours by the location of the pain, and by the swelling being in the iliac fossa rather than in the loin. In some cases, however, the pain may be referred to the lumbar region, or an appendiceal inflammatory mass may spread upwards. This is especially so when the appendix is retrocaecal in position. The onset of the trouble, the acute symptoms and the febrile disturbance will usually distinguish these cases from renal lesions.

Malignant growths of the large intestine
Malignant growths of the large intestine, especially of the ascending or descending colon, may form a mass in the loin that closely resembles a renal swelling. The mass formed by the growth may be grasped bimanually, is movable in the same directions as a renal tumour, and comes forward under the costal margin. The percussion note over the front of the lump is resonant, and there is usually an aching pain in the loin. If the growth has infiltrated through the wall of the bowel uncovered by peritoneum, the peri-renal tissues may be thickened, or proteinuria may be produced by direct invasion of the kidney, when the case will even more resemble a renal lesion. Carcinoma of the large intestine should be suspected if there is any irregularity in the action of the bowels, mucus or blood in the motions, or any symptom of incipient obstruction in the intestine. The tumour may be irregular and nodular, whereas a renal tumour presents rounded margins. The occurrence of a tumour in either side, associated with discomfort or palpable distension of the caecum from the accumulation of faeces, would render a growth in the colon the more suspicious.

The diagnosis of a large bowel tumour can usually be established by a barium enema X-ray examination or CT scanning with oral contrast. Confirmation can be made by direct colonoscopic examination, at which biopsy material can usually be obtained for histological examination.

Tumours of the omentum, mesentery or pancreas
These tumours, either cystic or malignant, are more median in position, do not project into the loin and seldom resemble a renal tumour. Retroperitoneal and peri-renal tumours may closely simulate renal tumours but will be distinguished on ultrasonography and certainly by CT scanning or magnetic resonance imaging.

THE DIFFERENTIAL DIAGNOSIS OF RADIOGRAPHIC SHADOWS IN THE ABDOMEN AND THE PELVIS
It is necessary for the true interpretation of radiographs that a clear conception should be held of the various conditions that may cast a shadow on an X-ray negative. In the diagnosis of cases of urinary disease, much information may be gained by the use of X-rays, and not merely to confirm the presence of calculi in some part of the urinary tract – on a good film, the outline and the size of the kidney can be seen. On a good film after efficient alimentary preparation, the outline of a normal kidney should be visible, lying opposite the bodies of the first, second and third lumbar vertebrae (Fig. K.7), and having an excursion of 4–5 cm in forced inspiration and expiration.
The following are the most frequent causes of a shadow that may be mistaken for a calculus:

- Intestinal contents including tablets
- Calcification of the mesenteric lymph nodes
- Gallstones, on the right side
- Calcification of the costal cartilages
- Caseous masses in a tuberculous kidney
- Areas of calcification in a renal growth
- Foreign bodies

*Intestinal contents* may cast a shadow in the renal area owing to inefficient preparation of the patient, or to the fact that he or she has recently taken bismuth, magnesium salts, etc. If any doubt exists, a second examination should be made. There may be some residue in the intestine from a recent barium-meal examination.

*Calcification of the abdominal or mesenteric lymph nodes* may cause a shadow in any part of the abdominal cavity. Although they are most frequently seen near the lower lumbar vertebrae or about the sacroiliac joint, and therefore external to the renal shadow, they may be superimposed upon the latter and cause difficulty in diagnosis. The shadow of a calcified node is usually mottled in appearance, small areas in the shadow showing increased density owing to the irregular deposition of lime salts; calcareous nodes are frequently multiple, but their chief characteristic is their range of mobility. A calcified node may be placed immediately in front of the kidney and move equally with it, causing great difficulty in diagnosis, or there may be a calculus in one kidney and calcareous nodes imitating calculi on the other side. An intravenous pyelogram or kidney–ureter–bladder (KUB) CT scan will usually clearly show the relation of a calculus to the renal pelvis.

*Gallstones* may give a shadow in the renal area on the right side. They are frequently multiple and may be seen to be faceted in a fusiform collection presenting the shape of the distended gallbladder (Fig. K.8). A single gallstone superimposed on the renal shadow may cause difficulty; the shadow of a gallstone is

**Figure K.7** Normal intravenous urogram at 30 minutes, following the removal of compression.

**Figure K.8** Intravenous urogram also showing gallbladder calculi and calcified mesenteric nodes, which can also be seen in the upper, plain, radiograph.
less dense than that of a renal stone and is frequently more dense in the central than in the peripheral part. In a lateral view, a stone in the gallbladder will occupy an anterior position in the abdomen, although one impacted in the common bile duct may be seen opposite the body of the first or second lumbar vertebra; in this case, there will probably be jaundice. In a cholecystographic examination, a gallstone may cause a filling defect (negative shadow) in the area of the gallbladder occupied by the dye. The distribution of stones in a horseshoe kidney may cause confusion until a pyelogram is done.

*Calcification of the costal cartilages* may give a shadow in the renal area in an anteroposterior negative. The shadows are not dense, are hazy in outline, and tend to assume a horizontal or oblique axis. On a lateral view, they will be placed immediately under the anterior abdominal wall.

With *calcified masses in a tuberculous kidney*, the shadow is rarely as defined as that of a calculus, is of moderate density with blurred and indistinct margins, appearing as one or more blotches in the renal area, but may occasionally, from the deposition of calcium salts, be very like the radiograph of a calculus (see Fig. H.5).

*Calcareaous areas in a renal carcinoma*: rarely, faint ill-defined areas may be present in a cystic (Bosniak types III and IV cysts) renal carcinoma. There may be symptoms of growth, such as haematuria and renal tumour, while a contrast CT examination will show a deformity of the pelvis and renal calices.

A foreign body, such as a shrapnel bullet, surgical clip or embolization coil, lying in front of or behind the kidney may mimic a calculus.

The line of the normal ureter lies anatomically along or just internal to the tips of the transverse processes of the second to fifth lumbar vertebrae, and it passes with a slight curve outwards in front of the sacroiliac articulation and then with a marked curve forwards and inwards to the base of the bladder. A shadow in this line may be due to a calculus in the ureter, but it must be differentiated carefully from other conditions. A calculus is usually small, rounded or oval, with a long axis in the line of the ureter. It may be found in any part of the course of the ureter, but it is seen most frequently in the lower end just before it enters the bladder. The conditions that may give a shadow that is likely to be mistaken for a ureteric calculus are:

- A calcified lymph node
- Calcification of the appendix or intestinal contents
- Phleboliths in the pelvis
- A foreign body

*Calcified lymph nodes* in the line of the ureter are placed most frequently in the angle between the last lumbar vertebra and the ala of the sacrum. They are usually multiple, forming a group in this situation in triangular form rather than in the longitudinal axis of the ureter; they are mottled in appearance, of irregular density, and are so movable that their position varies in successive radiographs.

A *calcified appendix* may occasionally give rise to a shadow in the line of the right ureter, suggesting a calculus, with very similar clinical symptoms. Further examination with a radio-opaque catheter in the ureter will show that the shadow is extrarenal.

*Phleboliths in the pelvis* are liable to be mistaken for ureteric calculi, but they often have a characteristic ring-like appearance, which is quite diagnostic. They are usually multiple and placed towards the peripheral areas of the pelvis, often about the level of the ischial spine. It must not be forgotten that a calculus may be present in the ureter in addition to phleboliths, but an IVU or KUB CT scan will differentiate them. Figure K.9 shows that the shadow of the ureter does not impinge on any of the numerous phleboliths present in the pelvis.

Figure K.9 Excretion pyelogram showing numerous phleboliths in the pelvis; the left ureter is seen passing between them. Note the 'bite defect' in the bladder on the right caused by a solid carcinoma.
Foreign bodies, especially shrapnel after periods of war, may occasionally lie near the line of the ureter. They are usually denser than calculi.

A shadow may be present in a pelvic radiograph that must be differentiated from that of a vesical calculus. The latter is usually rounded or oval, occupies a fairly central position in the pelvis, and may show rings of varying density, owing to the deposition of layers of urinary salts of different composition. Occasionally, one or more vesical calculi may form a shadow in a more lateral position in successive negatives, when a suspicion of their presence in a diverticulum in the bladder will arise. The diagnosis of this condition is discussed below. The following conditions may give rise to radiographic shadows in the pelvis:

- **Prostatic calculi:** these may be single or multiple, but in the radiograph they occupy a position very low in the pelvis, often behind the shadow of the pubis (Fig. K.10).
- **Calcification of a uterine fibroid:** this produces a large, irregular shadow of varying density. Bimanual palpation of a tumour moving with the uterus points to the diagnosis.
- **Opaque masses in a dermoid cyst of the ovary:** irregular shadows in the pelvis due to the formation of bone or teeth in the cyst. They may be present in young adult life, and a tumour would be palpated on abdominal or pelvic examination.
- **Calcification of a bladder tumour:** usually faint, ill-defined shadows on the pelvic X-ray. A cystoscopy reveals the true nature.
- **Foreign bodies in the bladder:** calcified through encrustation, or introduced by intent or iatrogenically (e.g. resectoscope tips or catheters).
- **Urethral calculus:** retained in the canal behind a stricture and enlarge in situ. They form a shadow in a radiograph above or below the pubic arch (Fig. K.11).

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*Figure K.10* Radiograph showing shadows in the pelvis due to multiple prostatic calculi.

*Figure K.11* Pelvic radiograph showing large calculus in the prostatic urethra.
LEG, ULCERATION OF

VENEROUS ULCER

Venous, or gravitational, ulcer almost invariably results from a preceding episode of deep vein thrombosis (DVT). There is usually a history of a painful swollen leg following childbirth, surgical operation or a period of recumbency from any disease, manifested by an episode of slight pyrexia, calf pain and tenderness and swelling of the leg as a result of thrombosis of the deep veins of the leg. Recanalization results in damage to the venous valves. Over subsequent years, the leg becomes swollen, superficial varicose veins may form due to perforator incompetence secondary to the raised venous pressure, pigmentation of the skin occurs, particularly over the medial side of the leg just above the medial malleolus (the pigment being haemosiderin), the subcutaneous, affected skin becomes eczematous, and the subcutaneous fat becomes replaced by thick fibrous tissue. Ulceration occurs as a consequence of the poor skin nutrition, either following some minor trauma or from scratching of the eczematous skin.

The patient may well have forgotten the original episode of deep vein thrombosis, perhaps many years before, and will blame the secondary varicose veins (as may, indeed, the medical practitioner) on the consequent ulcer, although it is unlikely that primary varicose veins alone give rise to this condition.

A venous ulcer is situated on the gaiter area of the leg, has a sloping edge and a floor that may be made up of red, velvety granulations if the ulcer is uninfected, white and fibrous if it is long-standing, and purulent and smelly if it is infected (Fig. L.1). If the ulcer is large, it has a sloping edge and a floor that may be made up of red, velvety granulations if the ulcer is uninfected, white and fibrous if it is long-standing, and purulent and smelly if it is infected (Fig. L.1). If the ulcer is large, the foot is often oedematous, so that the pedal pulses may not be palpable. It is important to exclude co-existing arterial disease (see below) and the presence or absence of these pulses must be determined with the Doppler probe.

ARTERIAL

Atherosclerosis and the other less common causes of arterial insufficiency may result in ulceration of the leg (see GANGRENE, PERIPHERAL, p. 216). This is particularly likely to occur over the pressure areas of the heel and malleolus rather than the gaiter area. There are the other features of peripheral arterial disease, including an absence of pulses, but it is important to remember that many elderly patients with venous ulceration of the leg have co-existing arterial disease, and this must be excluded. The use of the Doppler probe is particularly valuable in this respect (see above).

LEG, ULCERATION OF

Harold Ellis

Although it is true that the great majority of leg ulcers are due to venous disease – all too often called, and inaccurately labelled, ‘varicose ulcers’ – it must be remembered that a large number of other conditions can lead to this common clinical situation.

The various aetiologies of leg ulceration can be classified into:

- Venous – following deep vein thrombosis
- Arterial – especially arteriosclerosis (see GANGRENE, PERIPHERAL, p. 216)
- Mixed venous and arterial
- Neuropathic – especially diabetes mellitus
- Chronic infection (e.g. gumma and pyoderma gangrenosa)
- Complicating systemic disease (e.g. rheumatoid arthritis, haemolytic anaemias and ulcerative colitis).
- Trauma – including Munchausen’s syndrome
- Malignant disease – Marjolin’s ulcer and malignant melanoma

LACRIMATION

Reginald Daniel

Lacrimation or tearing is a function of the lacrimal gland and the accessory lacrimal glands. Tears wet the surface of the cornea and conjunctivae, thereby protecting the surface epithelium. They also inhibit the growth of organisms, provide the cornea with nutrient substances, and make the cornea a smooth optical surface by abolishing minute surface irregularities.

Basal tear production provides enough tears for these purposes, and reflex lacrimation occurs with irritative conditions or corneal disease (see EPIPHORA, p. 161). Psychic lacrimation is normally associated with pain or emotional upset.

Tear production diminishes with age, and dryness of the eyes is a common complaint of the elderly. Excessively dry eyes (xerophthalmia) can result idio pathically, congenitally and in cases of autoimmune disorders including rheumatoid arthritis and Sjögren’s syndrome. The dry eye condition can follow trachoma infections, the use of certain drugs including oral contraceptives and following refractive surgery. The patient complains of a gritty or burning sensation in the eyes. Artificial tears used frequently are usually beneficial, and surgical occlusion of the nasolacrimal drainage system can be performed in severe cases.

LEG, ULCERATION OF

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- Malignant disease – Marjolin’s ulcer and malignant melanoma
NEUROPATHIC
Loss of sensation in the lower limb may result in trophic ulceration as a result of the pressure of the shoe, or an orthopaedic appliance, etc. Both in the industrialized world and increasingly in the Asian subcontinent, the most common cause of neuropathic ulcers is diabetes. Typical sites are over the malleoli, on the heel or on the sole of the foot over the heads of the metatarsals. In the severe diabetic, this neuropathy may be complicated by the associated vascular impairment of diabetic microangiopathy and by the increased propensity of the diabetic to infection (Fig. L.2).

Other examples of neuropathic ulceration will be seen in conditions such as leprosy, tabes dorsalis and hemiplegia following a stroke.

CHRONIC INFECTION
Syphilitic ulcers are the result of breakdown of a gumma that has formed in the subcutaneous tissues in tertiary syphilis. These ulcers occur in the upper third of the leg (in contrast to venous ulcers, which are confined to the lower third). They are almost always circular and present a punched-out appearance. They are generally multiple and tend to run into each other, forming a large ulcer with a serpiginous outline. There may be other signs of syphilis, and there will be positive serological tests for this disease. However, today it is rarely seen in the Western world.

*Pyoderma gangrenosum* produces multiple ulcers, which may occur anywhere in the body but frequently affect the leg. They are multiple and have ragged blue–red overhanging edges and necrotic bases. They often start as tender infected nodules, often following minor trauma, and are usually associated with patients suffering from chronic ulcerative colitis or Crohn’s disease.

Leg ulcers may be seen in amoebiasis, chancroid, diphtheria, leprosy, yaws and kala-azar, all of which might be considered in recent immigrants. Chronic sores and ulcers of the leg, as with other parts of the body, may be due to various skin fungi (e.g. blastomycosis, sporotrichosis and actinomycosis).

Figure L.1 Varicose (venous) ulcer. Note the extensive surrounding pigmentation due to deposition of haemosiderin.

Figure L.2 (a, b) Diabetic leg ulcers due to peripheral neuropathy.
Multiple discharging sinuses in a swollen and distorted foot may be seen in Madura foot, resulting from infection with the fungus *Nocardia madurae*.

Lupus vulgaris, a form of primary tuberculosis of the skin, is not often found on the leg, although it may occur there as in any cutaneous area.

**SYSTEMIC DISEASE**

Ulcers may complicate a large variety of systemic diseases. Pyoderma gangrenosum in ulcerative colitis and Crohn's disease has already been noted. Ulcers of the leg may also occur in rheumatoid arthritis, sickle-cell anaemia, thalassaemia, polycythaemia rubra vera, thrombocytopenic purpura, hereditary spherocytosis and the leukaemias. All of these might be considered where the aetiology is not evident.

**TRAUMA**

Trauma, unless it is continuous, is not alone sufficient to cause an ulcer in healthy people. Old ladies, with thin atrophic skin, may lacerate the tissues over the shin, and the poor blood supply may result in necrosis and subsequent ulceration of the damaged skin. Ulceration may result in inadvertent permeation of the subcutaneous tissues by the sclerosing fluid used in the injection treatment of varicose veins. The malingerer may rub corrosive agents into the leg or use a coin bandaged firmly against the skin (examples of the so-called Munchausen's syndrome). The diagnosis is often suggested by the rectangular or other definite shape of the ulcer itself, as well as by the strange personality of the patient.

**MALIGNANT DISEASE**

A carcinoma may develop in a chronic ulcer, particularly a venous ulcer that has existed for many years (Marjolin's ulcer). The ulcer at one edge becomes heaped up, everted and indurated (Fig. L.3). The inguinal lymph nodes may become enlarged. The appearance of bare bone at the base of a venous ulcer should always arouse the greatest suspicion that malignant change has taken place. A biopsy specimen at the edge of the ulcer should be removed for histological examination in any case of doubt.

A malignant melanoma may ulcerate and bleed (Fig. L.4). Soft tissue or bone tumours may, at a late stage, fungate through the skin and give rise to an irregular breaking-down mass.

**LIBIDO, EXCESSIVE**

Paul Carroll

The desire for sexual intercourse in humans is a biological variable that ranges from the diminutive to the excessive. Sexual interest may be greater than desirable, as reflected in the terms ‘hypersexuality’, ‘sex addiction’ and ‘satyriasis’. At what point the sex drive becomes excessive is not easy to determine,
since the normal sex drive can be considerable, but if the individual feels distress as a consequence, this can be a reason to regard the libido as excessive. The causes of excessive libido include conditions associated with androgen excess, psychological health issues and medications, and are listed in Table L.1. Females may show excessive libido during the manic phase of a manic–depressive (bipolar) psychosis. Precocious puberty in a boy (see PUBERTY, PRECOCIOUS, p. 535) or a girl may lead to a libido that is excessive for that individual, in that it is inappropriate for the age. A similar situation may be found in untreated congenital adrenal hyperplasia, commonly due to a 21-hydroxylase deficiency, when there is a build-up of androgens from hypersecretion by the adrenal gland. The result is raised testosterone levels in boys and girls, with penile and clitoral enlargement. Since androgens are important in maintaining libido in women, any situation in which a woman is exposed to excess androgens may promote excessive libido. These include androgen administration, polycystic ovary syndrome (see HIRSUTISM, p. 281) or the very rare case of androgen-secreting tumour of the adrenal gland or ovary. Most infections tend to reduce libido, but, curiously, it was noticed in both sexes, at a time when chronic tuberculosis was rife, that patients were prone to an increase in their sexuality. Whether this was a true effect of the disease is unknown. Alcohol may stimulate libido in both sexes in small doses, but larger amounts, particularly if taken chronically, tend to depress libido. Some cases of increased libido in women have been described following opioid or benzodiazepine withdrawal.

**LIBIDO, LOSS OF**

Paul Carroll

Libido relates to sexual desire and drive and is influenced by both psychological and hormonal factors in both genders. Circulating androgens are promoters of sex drive and are derived largely from the testes in men and from the adrenal glands in women. Thus, hypothalamic or pituitary disease, which reduces gonadotrophin levels in men or adrenocorticotropic hormone levels in women, will deprive each sex respectively of their main source of androgen. Similarly, testicular disease or damage in a male or primary adrenal failure (e.g. Addison's disease) in a female will achieve the same effect.

A reduction in gonadotrophins may occur in chronic alcoholics. In addition, cirrhosis of the liver gives rise to an increased oestrogen:testosterone ratio in a male, which may also lead to a reduction in libido. An increased oestrogen: testosterone ratio is also found in thyrotoxicosis. Oestrogen treatment or downregulation of gonadotrophins by long-acting gonadotrophin-releasing hormone analogues in the treatment of carcinoma of the prostate lead to loss of libido. Hyperprolactinaemia may give rise to a reduction in libido in men by reducing the secretion of gonadotrophins, and by blocking the action of gonadotrophins on the testes. The cause of hyperprolactinaemia may be a pituitary tumour (prolactinoma) or other hypothalamic and pituitary disease leading to a lack of prolactin-inhibiting factor (dopaminergic tone). Raised prolactin levels may also be found in primary hypothyroidism and renal failure, and as a result of drug therapy (Table L.2). Anti-androgens used in the treatment of benign prostatic hypertrophy or carcinoma may lead to loss of libido.

Psychological problems are important as causes of loss of libido. Some people just have naturally low levels of libido. Others are anxious or depressed. The effect of childhood trauma on future sexual behaviour has not been fully evaluated. Local causes of dyspareunia such as a median episiotomy scar, previous pelvic surgery and a relaxed vaginal outlet secondary to multiple vaginal deliveries can give rise to reduced sexual enjoyment for women. Diminished libido has been documented in women taking

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**Table L.1 Causes of excessive libido**

<table>
<thead>
<tr>
<th>Male</th>
<th>Female</th>
</tr>
</thead>
<tbody>
<tr>
<td>Constitutional</td>
<td>Constitutional</td>
</tr>
<tr>
<td>Psychological</td>
<td>Psychological</td>
</tr>
<tr>
<td>Manic phase of manic–depressive (bipolar) psychosis</td>
<td>Menopause</td>
</tr>
<tr>
<td>Precocious puberty</td>
<td>Psychological</td>
</tr>
<tr>
<td>Congenital adrenal hyperplasia</td>
<td>Psychological</td>
</tr>
<tr>
<td>Androgen-secreting tumour of suprarenal gland or ovary</td>
<td>Androgen therapy</td>
</tr>
<tr>
<td>Infections</td>
<td>Infected</td>
</tr>
<tr>
<td>Tuberculosis</td>
<td>Tuberculosis</td>
</tr>
<tr>
<td>Drugs</td>
<td>Drugs</td>
</tr>
<tr>
<td>Alcohol (in small doses)</td>
<td>Alcohol (in small doses)</td>
</tr>
<tr>
<td>Following opioid or benzodiazepine withdrawal</td>
<td>Following opioid or benzodiazepine withdrawal</td>
</tr>
</tbody>
</table>

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Alcohol may stimulate libido in both sexes in small doses, but larger amounts, particularly if taken chronically, tend to depress libido. Some cases of increased libido in women have been described following opioid or benzodiazepine withdrawal.
### Table L.2 Causes of loss of libido

<table>
<thead>
<tr>
<th>Condition</th>
<th>Mechanism</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Male</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Endocrine</strong></td>
<td></td>
</tr>
<tr>
<td>Hypothalamic disease</td>
<td>Low gonadotrophins and thyroid-stimulating hormone</td>
</tr>
<tr>
<td>Pituitary disease</td>
<td>Pituitary disease</td>
</tr>
<tr>
<td>Thyrotoxicosis</td>
<td>Raised oestrogen:testosterone ratio</td>
</tr>
<tr>
<td>Hypothyroidism</td>
<td>Psychomotor retardation, lethargy, raised prolactin</td>
</tr>
<tr>
<td>Cushing’s disease</td>
<td>Decreased luteinizing hormone</td>
</tr>
<tr>
<td>Feminizing tumour of testis or suprarenal gland</td>
<td>Oestrogen secretion</td>
</tr>
<tr>
<td>Testicular disease or castration</td>
<td>Reduced testosterone</td>
</tr>
<tr>
<td>Ageing</td>
<td>Reduced testosterone</td>
</tr>
<tr>
<td><strong>General disease</strong></td>
<td></td>
</tr>
<tr>
<td>General debilitating disease (e.g. chronic infection, cancer)</td>
<td>Psychological</td>
</tr>
<tr>
<td>Chronic alcoholism</td>
<td>Reduced gonadotrophins</td>
</tr>
<tr>
<td>Cirrhosis of the liver</td>
<td>Raised oestrogen:testosterone ratio</td>
</tr>
<tr>
<td>Renal failure</td>
<td>General debility</td>
</tr>
<tr>
<td><strong>Psychological</strong></td>
<td></td>
</tr>
<tr>
<td>Idiopathic lack of sexual desire</td>
<td></td>
</tr>
<tr>
<td>Depression</td>
<td></td>
</tr>
<tr>
<td>Anxiety</td>
<td></td>
</tr>
<tr>
<td><strong>Drugs</strong></td>
<td></td>
</tr>
<tr>
<td>Gonadotrophin-releasing hormone analogues</td>
<td>Reduced gonadotrophins</td>
</tr>
<tr>
<td>Oestrogens</td>
<td>Testicular damage</td>
</tr>
<tr>
<td>Cytotoxic agents</td>
<td></td>
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<tr>
<td>Alcohol</td>
<td></td>
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<tr>
<td>Phenothiazines</td>
<td></td>
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<tr>
<td>Metoclopramide</td>
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<tr>
<td>Haloperidol</td>
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<td>Pimozide</td>
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<tr>
<td>Methyldopa</td>
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<tr>
<td>Reserpine</td>
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<tr>
<td>Cimetidine</td>
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<tr>
<td>Spironolactone</td>
<td></td>
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<tr>
<td>Cimetidine</td>
<td></td>
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<tr>
<td>Cyproterone acetate</td>
<td></td>
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<tr>
<td>Flutamide</td>
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<tr>
<td>Finasteride</td>
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<tr>
<td><strong>Female</strong></td>
<td></td>
</tr>
<tr>
<td><strong>Endocrine</strong></td>
<td></td>
</tr>
<tr>
<td>Hypothalamic disease</td>
<td>Reduced adrenocorticotrophic hormone</td>
</tr>
<tr>
<td>Pituitary disease</td>
<td>Psychomotor retardation, lethargy</td>
</tr>
<tr>
<td>Hypothyroidism</td>
<td>Decreased secretion of androgens</td>
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<tr>
<td>Addison’s disease</td>
<td>Decreased adrenal androgens</td>
</tr>
<tr>
<td><strong>Psychological</strong></td>
<td></td>
</tr>
<tr>
<td>Depression</td>
<td></td>
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<tr>
<td>Anxiety</td>
<td></td>
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<tr>
<td><strong>Drugs</strong></td>
<td></td>
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<tr>
<td>Cyproterone acetate</td>
<td>Anti-androgens</td>
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<tr>
<td>Alpha-and beta-blockers</td>
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<tr>
<td>Opiates</td>
<td></td>
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<tr>
<td>Chronic alcohol excess</td>
<td></td>
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<tr>
<td><strong>Systemic disease</strong></td>
<td></td>
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<tr>
<td>Pelvic and breast neoplasms</td>
<td>Raised prolactin</td>
</tr>
<tr>
<td>Myocardial infarction</td>
<td></td>
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<tr>
<td>Renal failure</td>
<td></td>
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<tr>
<td>Irritable bowel syndrome</td>
<td></td>
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<tr>
<td><strong>Infections</strong></td>
<td></td>
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<tr>
<td>Vaginitis:</td>
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<tr>
<td>Trichomonas</td>
<td></td>
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<tr>
<td>Yeasts</td>
<td></td>
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<tr>
<td>Bacteria</td>
<td></td>
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<tr>
<td>Pelvic inflammatory disease</td>
<td></td>
</tr>
<tr>
<td>HIV</td>
<td>Advanced AIDS giving rise to ovarian failure</td>
</tr>
<tr>
<td><strong>Local</strong></td>
<td></td>
</tr>
<tr>
<td>Dyspareunia</td>
<td></td>
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<tr>
<td>Endometriosis</td>
<td></td>
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<tr>
<td>Menopause</td>
<td></td>
</tr>
<tr>
<td>Vaginal dryness secondary to oestrogen deficiency</td>
<td>Atrophy of urogenital tissue</td>
</tr>
<tr>
<td>Episiotomy scarring</td>
<td></td>
</tr>
<tr>
<td>Post-radiation vaginal atrophy</td>
<td></td>
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<tr>
<td>Vulvitis</td>
<td></td>
</tr>
<tr>
<td>Infection of Bartholin’s gland</td>
<td></td>
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<tr>
<td>Hymenal obstruction</td>
<td></td>
</tr>
<tr>
<td>Pelvic surgery</td>
<td></td>
</tr>
<tr>
<td>Relaxed vaginal musculature</td>
<td></td>
</tr>
<tr>
<td>Reduced sexual enjoyment secondary to multiple pregnancies</td>
<td></td>
</tr>
</tbody>
</table>

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alpha- and beta-blockers, opiates, anti-androgen medications and some psychiatric drugs.

**LIPS, AFFECTIONS OF**

*Mark Kiniron*

The lips form an important mucocutaneous junction, and their proper function is important for feeding and communication. They are subserved by a very large quantity of nerves, and hence inflammation can cause disproportionate irritation and pain. The lips are also of considerable cosmetic, psychological and emotional importance (Box L.1).

Macular affections (see MACULES p. 396) around the lips include flat moles, freckles and the multiple tiny dark freckles seen in Peutz–Jegher’s syndrome. Multiple lip telangiectases are seen in hereditary haemorrhagic telangiectasia (see TONGUE, DISCOLORATION OF, p. 685) and sometimes with severe liver disease.

The most common papules on lips are probably plane warts, but lichen planus papules have a predilection for the lips, as do plaques of discoid lupus erythematosus. Pyogenic granuloma can form at this site, and in older patients venous lakes are commonly seen. The lip can be the site of a primary syphilitic chancre. Actinic cheilitis is common on the lower lip, especially in seafarers and agricultural workers. The appearance is of atrophy with greyish plaques often surmounted by crusts in cold weather. Solar keratoses may occur on the lips in early, as metastases can quickly occur. Molluscum contagiosum and viral warts may occur on the lips in people with immunosuppression from HIV infection. Swollen lips can follow trauma and thermal insult, or be part of an urticaria, as angio-oedema (see WEALS, p. 754). The Melkerson–Rosenthal syndrome comprises permanently oedematous thickened lips (granulomatous cheilitis) with recurrent facial palsy and scrotal tongue. Granulomatous cheilitis is occasionally seen in Crohn’s disease. Thickening of the skin around the mouth (rather than swelling) is a characteristic feature of atopic dermatitis due to rubbing of the lips with the backs of the hands.

Erosions on the lips occur in acute infections such as impetigo in children, herpes simplex, which may be recurrent in adults, hand-foot-and-mouth disease and secondary syphilis. Erosive dermatoses that affect the lips include erythema multiforme (Stevens–Johnson syndrome; see p. 426) and fixed drug eruption (codeine and sulphonamides). Cocaine smoking is a direct cause of lip erosions. Chronic erosions around the lips are seen in zinc deficiency, acrodermatitis enteropathica and pemphigus (see p. 425).

When inflammation is confined to the lips, a search should be made for occult candidiasis, especially in those with dentures and iron deficiency. Exfoliative cheilitis may occur in patients with HIV infection and oral candidiasis. Chronic contact dermatitis can occur at this site – for example, nickel dermatitis from sucking hairpins, lipstick dermatitis, toothpaste dermatitis and, rather surprisingly, nail varnish dermatitis. Irritation from excessive lip-licking can often be observed in patients with cheilitis and may sometimes be the primary cause.

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**Box L.1** Affections of the lips

<table>
<thead>
<tr>
<th>Macules</th>
<th>Melkerson-Rosenthal syndrome</th>
<th>Crohn’s disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flat mole (junctional naevus)</td>
<td>Impetigo</td>
<td>Erosions</td>
</tr>
<tr>
<td>Peutz–Jegher’s syndrome</td>
<td>Herpes simplex</td>
<td></td>
</tr>
<tr>
<td>Telangiectasia</td>
<td>Hand-foot-and-mouth disease</td>
<td></td>
</tr>
<tr>
<td>Hereditary haemorrhagic telangiectasia</td>
<td>Secondary syphilis</td>
<td></td>
</tr>
<tr>
<td>Liver disease</td>
<td>Erythema multiforme</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Fixed drug eruption</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Zinc deficiency</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Acrodermatitis enteropathica</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Pemphigus</td>
<td></td>
</tr>
<tr>
<td>Erosions</td>
<td>Thinning</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Atopic dermatitis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lip-licking</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cheilitis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Candidiasis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Contact dermatitis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Lip-licking</td>
<td></td>
</tr>
</tbody>
</table>

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**LIVER, ENLARGEMENT OF**

*Simon Anderson*

The normal liver can be palpable in children, in thin people with lax abdominal muscles, and in those with chronic obstructive airways disease. Palpation of the liver is an unreliable indicator of actual liver size, and percussion can be more accurate. Hepatic dullness extends from the fifth intercostal space in the right nipple line to the seventh intercostal space in the mid-axillary line with a span of about 12 cm.

In health, the edge of the liver is firm and uniform, and the surface feels smooth. A tongue-like projection of the right lobe may protrude from its lower right-hand part. This projection, known as Riedel’s lobe, is more common in women than in men. It may cause difficulty of diagnosis, being confused with a mobile kidney, gallbladder or tumour.
Many conditions that are unconnected with the liver cause an apparent alteration in its size. In emphysema, the liver is easily palpable, but percussion will reveal that the organ is merely displaced downwards by the hyperinflated lungs. Deformities of the chest due to rickets or curvature of the spine may depress the liver, as may a right subphrenic abscess. It is unusual for enlargement of the liver to lead to upward extension of hepatic dullness because the weight of the liver causes it to descend. Elevation of the upper limit of hepatic dullness occurs when local hepatic disease involves the diaphragm (amoebic abscess or hydatid cyst). Loss of hepatic dullness occurs in emphysema, principally because of displacement of the liver. Free gas in the peritoneum or distension of the colon may also do this.

Hepatoptosis and wandering liver are terms applied to a liver that is found in an abnormal position. This is rare, but it does occur after therapeutic pneumoperitoneum at laparoscopy. It is usually an asymptomatic condition, although the patient may complain of a dragging sensation and heaviness in the right upper quadrant of the abdomen. The liver that is displaced may be thought to be enlarged.

Hepatomegaly can be a feature of any acute hepatitis whether infectious (hepatitis A, B, C and E, cytomegalovirus, Epstein–Barr virus, herpes simplex or toxoplasma), toxic (alcohol, medications), metabolic (haemachromatosis, Wilson’s disease and fatty liver of pregnancy) or autoimmune disease. Such hepatic enlargement is often tender and associated with jaundice and sometimes fever.

Vascular causes of acute hepatomegaly are uncommon. Hepatomegaly and ascites are features of Budd–Chiari syndrome, thrombosis of the hepatic veins that may occur due to a hypercoagulable state (hormonal medication, malignancy or genetic clotting abnormalities) or a vascular web at the junction of the hepatic vein and inferior vena cava. Venous congestion secondary to right heart failure, tricuspid incompetence or constrictive pericarditis may result in tender, pulsatile hepatomegaly with peripheral oedema, ascites, jaundice and markedly raised jugular venous pressure. Veno-occlusive disease of the liver is an increasingly recognized condition that is caused by ingestion of plants in the form of drinks (‘bush-tea’) and chewing (kava). Sickle-cell crisis and malaria-associated haemolysis are further causes of tender hepatomegaly. Infiltration by cancer (non-Hodgkin’s lymphoma, metastases) or primary hepatocellular cancer in patients with pre-existing cirrhosis due to chronic viral hepatitis (hepatitis B or C virus), alcohol or haemochromatosis is often tender. The co-existence of splenomegaly or the detection of lymphadenopathy may indicate lymphoma. Metastatic neuroendocrine gut tumours (formerly known as carcinoid tumours) are associated with the symptoms of the carcinoid syndrome, namely facial flushing, diarrhoea, and occasionally asthma and valvular heart disease (pulmonary stenosis).

Infiltrative conditions such as sarcoid, amyloid and fatty liver tend to give rise to non-tender hepatomegaly. So-called ‘fatty liver’ (steatosis) is a common condition found particularly in those with type 2 diabetes, hyperlipidaemia and obesity (called non-alcohol fatty liver disease). It can also occur after jejunoo-ileal bypass and, paradoxically, protein malnutrition. Although usually benign, it can give rise to significant hepatitis, called non-alcoholic steatohepatitis (NASH), which sometimes result in cirrhosis. An acute form of this condition is fatty liver of pregnancy, which occurs in the third trimester and may be fatal. Hepatic sarcoid is of little clinical consequence to the liver, but it may indicate other visceral involvement such as myocardial infiltration, particularly in African-Caribbean Americans, and carries a poor prognosis.

In addition to viral disease, the liver is also a target for other infections such as tuberculosis, schistosomiasis, malaria, kala-azar, hydatid disease (Fig. L.5), liver flukes and Ascars lumbricoides (biliary ducts) and syphilis. In most of these conditions, the hepatomegaly is firm and non-tender, while with syphilis it is nodular. Acute painful hepatomegaly occurs during the crises of malaria. With schistosomiasis, pre-sinusoidal portal hypertension causes splenomegaly, and oesophageal varices then develop. Hepatic synthetic function is maintained until a late stage, and jaundice is relatively mild.

Figure L.5 Multiple cysts of the liver due to hydatid disease.
Acute bacterial infections and abscesses may originate from the portal vein (from a gastrointestinal source such as the appendix or diverticulitis), hepatic artery (systemic infection, e.g. endocarditis) or from the biliary tree (e.g. primary sclerosing cholangitis).

An amoebic abscess may be single and large. It commonly follows a history of dysentery, and the majority occur in the right lobe. Men are affected more often than women. The patient presents with swinging pyrexia, associated with rigors and tachycardia, and a considerably enlarged and very tender liver, and there may also be a sympathetic pleural effusion. These abscesses can be slow to reveal themselves and often require repeated ultrasound scans to be diagnosed.

Disseminated visceral *Candida* affecting the liver is a well-recognized complication of the early post-chemotherapy period when granulocyte counts are recovering.

Large hydatid cysts can occur within the liver, but these are usually asymptomatic and cause no disturbance of liver function. The cysts are rounded and smooth and may achieve considerable size. Calcified cyst can be seen on plain abdominal X-ray. Ultrasound appearances are characteristic, with ‘daughter’ cysts and internal septae often seen. CT scan can provide more detailed information. Needle aspiration is contraindicated because this may cause infection and spillage of cysts within the peritoneal cavity. Serum antibody ELISA (enzyme-linked immunosorbent assay) tests can confirm the diagnosis.

Simple cysts of the liver are very common and usually go undetected. They can be associated with cysts of the kidney and pancreas (polycystic kidney disease) and von Hippel–Lindau syndrome. Giant cysts can cause significant chronic symptoms due to their size, or rarely acute pain due to internal haemorrhage. Aspiration is of no value in treating these cysts, as they recur.

Hepatic haemangiomas are common and are generally of no clinical consequence other than for causing confusion when interpreting scans. Both fibrolamellar tumours and hepatic adenomas are well recognized; adenomas are associated with pregnancy and the use of oestrogens.

A variety of metabolic, genetic and endocrine disorders are associated with the development of hepatomegaly. *Glycogen storage diseases* cause hepatomegaly from birth. Lipid storage disorders such as *Gaucher’s disease* are associated with massive hepatosplenomegaly and growth retardation, while *Niemann–Pick disease* may have neurological associations, including mental retardation.

*Haemochromatosis* is an autosomal recessive disorder of iron metabolism. The disease is rare in premenopausal women because of menstrual blood loss. The liver is firm and often enlarged. There may be evidence of portal hypertension including splenomegaly and ascites, while a proportion of patients present for the first time with a variceal haemorrhage. Iron is also deposited in the skin, producing a dusky, slate-grey pigmentation, heart (dilated cardiomyopathy) and joints (arthropathy).

Deposition in the pancreas accounts for the common association with diabetes (‘bronze diabetes’) and involvement of other endocrine glands (pituitary and adrenals) leads to deficiency states. Marked feminization manifests as gynaecomastia, while absent body hair, testicular atrophy and a decreased need for shaving are prominent features. Although relatively unusual, the diagnosis of haemochromatosis is important because the prognosis is greatly improved by venesection. Massive hepatomegaly associated with haemochromatosis may be due to the development of hepatocellular cancer, which is a relatively common late complication.

Wilson’s disease is an autosomal recessive condition leading to the accumulation of copper, particularly in the liver (producing acute or chronic liver disease) and brain (producing neuro-psychiatric symptoms). The golden rings of copper deposited in Descemet’s membrane of the eyes seen on slit-lamp examination (Kayser–Fleischer rings) are diagnostic of central nervous system involvement. Blue lunulae may be seen on examination of the hands.

Acromegaly is associated with hepatomegaly, without evidence of liver dysfunction. The liver is modestly enlarged, but soft. Thyrotoxicosis may also be associated with hepatic enlargement and deranged liver function tests.

**DIAGNOSTIC INVESTIGATIONS**

The investigation of hepatomegaly usually involves an ultrasound scan with additional information being contributed by computed tomography (CT) and magnetic resonance imaging (MRI).

In cases of hepatomegaly, the liver blood tests may be helpful, with most causes giving a slight rise in alkaline phosphatase and possibly gamma-glutamyl transaminase. Serum tumour markers such as carcinoembryonic antigen (colorectal cancer), alphafetoprotein (germ cell tumours and hepatocellular carcinoma) and CA19.9 (pancreaticobiliary cancers) may provide some indication of the primary site of a supposed metastatic cancer. Serum angiotensin-converting enzyme activity is raised in sarcoidosis. Specific antibody tests are available to identify infectious agents.
Haemochromatosis is diagnosed with the finding of a raised serum ferritin and transferrin saturation, and confirmed with genotype testing (C282Y;H63D mutation). CT and MRI assessment of iron stores have been used in specialized centres.

Wilson's disease is diagnosed on the basis of a low serum ceruloplasmin level (carrier protein for copper), excessive urinary copper excretion and the presence of Kayser–Fleischer rings.

Neuroendocrine tumours giving rise to the carcinoid syndrome can be detected by urinary measurement of 5-hydroxyindoleacetic acid.

An ultrasound-guided liver biopsy is a generally safe procedure if the platelet count and clotting profile are normal. If a coagulopathy is present, a transjugular procedure if the platelet count and clotting profile are normal. If a coagulopathy is present, a transjugular procedure if the platelet count and clotting profile are normal. If a coagulopathy is present, a transjugular procedure if the platelet count and clotting profile are normal.

5-hydroxyindoleacetic acid.

syndrome can be detected by urinary measurement of Wilson's disease is diagnosed on the basis of a low serum ceruloplasmin level (carrier protein for copper), excessive urinary copper excretion and the presence of Kayser–Fleischer rings.

While a regional (anatomical) approach is very useful in diagnosing mechanical problems, do not overlook the challenge posed by dual pathology especially generalised diseases. For example, attributing a patient's problem with walking to obvious bow medial compartment osteoarthritis of the knee when the real cause is early motor neurone disease, is a classical error. Taking a proper history, examining the neurology, checking the peripheral pulses and observing the patient walk is the best defence against error.

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LEG PAIN OF RADICULAR OR VASCULAR ORIGIN
This section deals specifically with the causes of pain referred into the limb arising from local lesions rather than generalized diseases. The subdivision of the section is largely anatomical, but nerve root compression (‘sciatica’), ischaemia (intermittent claudication) and spinal stenosis (spinal claudication), which are responsible for a large proportion of pain in the buttock and leg, are considered separately first.

Sciatica
Sciatica is a term hallowed by common usage, both in the lay population and by doctors, but unfortunately it means a multiplicity of different things, and is therefore probably a term to be avoided. Asking the question ‘Is there any evidence that the patient has nerve root compression?’ is much more valuable. If the term is used, it should be restricted to those cases in which there is definite evidence of involvement of the sciatic nerve or one of its component roots. For example, pain which radiates from the lower back to the buttock and down the back of the thigh to the knee is quite commonly found in patients with a spondylolisthesis, who do not show any evidence of nerve root compression. Restrict ‘sciatica’ to a pain that radiates from the back down through the buttock, down the back of the thigh and down to the ankle or foot, passing along either the lateral or posterior aspect of the calf and aggravated by coughing or sneezing, together with the presence of nerve root tension signs, for example a positive straight leg raising test. Objective motor signs, such as loss of the ankle jerk (S1) or weakness of extension of the big toe (L5; extensor hallucis longus), are much more reliable in defining which nerve root is involved than sensory loss. Unfortunately for the clinician – but fortunately for the patient – radicular pain of recent onset is usually not accompanied by any objective neurological signs.

While lumbar degenerative disc disease and spondylosis account for the vast majority of cases of backache and referred pain into the legs, other lesions can be associated with pain on coughing, a positive Lasègue's straight leg raising test and a positive l'hermitte's sign – a surge of paraesthesia down the trunk into the limbs on forward flexion of the neck. Examples include multiple sclerosis, subacute combined degeneration of the cord, a cervical spinal tumour and cervical spondylosis causing cord compression.

The most common cause of radicular referred pain is a posterolateral disc protrusion at the L4/L5 or L5/S1 level. Classic disc disease is episodic, and often first presents in early adult life, being frequently precipitated by a ‘minor’ injury such as lifting a heavy weight. With recurrent episodes, the pain radiates further down the leg and the nerve root tension signs become more evident. However, there may be no such history, and symptoms from nerve root compression may be present from the first episode.
The lumbar lordosis may be lost or there may be a scoliosis, which becomes more marked on forward flexion. Tenderness is often – but by no means invariably – present at the level of the lesion, and also in the region of the posterior superior iliac spine. A characteristic of mechanical backache is that a good (if not full) range is retained in at least one direction. Areas of reduced sensation to pinprick, weakness and reflex changes associated with individual root lesions are summarized in Figure L.6 and Table L.3.

Beware of the patient who presents with referred pain into both legs or pain that alternates between the two legs as there may be a compression of the cauda equina. In particular, look for loss of sensation in the perianal area, and a loss of anal tone. Fortunately, cauda equina lesions due to a central disc prolapse are uncommon, but the symptoms can be vague and are frequently missed, especially when due to a slowly expanding benign tumour such as a neurofibroma.

Plain X-rays of the lumbar spine are usually unhelpful in diagnosing degenerative causes of symptoms as a young adult with a disc prolapse usually has normal X-rays, whereas from middle age onwards almost everybody shows some evidence of degenerative disease of the spine. The great significance of plain X-rays is that they help to exclude other more serious diseases, for example pyogenic and tubercular infection (‘involvement of two bodies and the intervening disc’); metastases (loss of a pedicle); collapse of a vertebral body due to either malignancy or an osteoporotic fracture; and benign tumours (a smooth erosion of one or more pedicles). Recent advances in imaging by computed tomography (CT) or magnetic resonance imaging (MRI) scanning have greatly increased diagnostic accuracy. To be significant, there must, particularly with degenerative lesions of the spine, be a match between the abnormality shown and the patient’s signs.

Box L.2 lists the causes of radicular pain radiating into the lower limbs. For further discussion on the following entities, see the appropriate cross-references:

- Intervertebral disc disease (see BACK, PAIN IN, p. 45).
- Spinal stenosis (see BACK, PAIN IN, p. 48).
- Spondylolisthesis (see SPINE, DEFORMITY OF, p. 627).
- Infection (see SPINE, DEFORMITY OF, pp. 628, 629 and Figs S.41 and S.42; SPINE, TENDERNESS OF, p. 633 and Figs S.51 and S.52).

Causes of intermittent and neurogenic claudication

These include:

- Spinal stenosis
- Leriche’s syndrome
- Iliac, femoral or popliteal stenosis
- Anaemia

---

**Figure L.6** Areas of reduced sensation to pinprick in lumbosacral root lesions (shaded, lumbar roots; white, sacral roots).

**Table L.3** Signs associated with common nerve root lesions affecting the legs

<table>
<thead>
<tr>
<th>Root</th>
<th>Paraesthesiae/numbness</th>
<th>Muscle weakness</th>
<th>Reflex change</th>
</tr>
</thead>
<tbody>
<tr>
<td>L1</td>
<td>Groin</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>L2</td>
<td>Front of mid thigh</td>
<td>Quadriceps</td>
<td>–</td>
</tr>
<tr>
<td>L3</td>
<td>Front of lower thigh</td>
<td>Quadriceps</td>
<td>Knee ↓</td>
</tr>
<tr>
<td>L4</td>
<td>Front of lower thigh, knee and inner aspect of shin</td>
<td>Quadriceps and tibialis anterior</td>
<td>Knee ↓</td>
</tr>
<tr>
<td>L5</td>
<td>Back of thigh, lateral aspect of leg, dorsum of foot to big toe</td>
<td>Extensor hallucis longus</td>
<td>–</td>
</tr>
<tr>
<td>S1</td>
<td>Back of leg, lateral aspect of foot and sole</td>
<td>Calf wasting and weakness of plantar flexors</td>
<td>Ankle ↓</td>
</tr>
</tbody>
</table>

N.B. It is uncommon to lose a knee jerk due to the dual innervation from L3 and L4.
The term **intermittent claudication** is used to describe the development of pains in the leg (usually the calf) that are brought on during walking, and which necessitate limping or rest. The pain is relieved after 5–15 minutes by rest, but it recurs after walking a similar distance. With the passage of time, the distance walked before pain becomes intolerable usually decreases, progressively restricting exercise tolerance. The condition is due to inadequate arterial blood supply to the leg muscles, and it is caused by narrowing of the iliac, femoral or popliteal arteries. Atheroma with or without thrombosis is the usual culprit, but embolism, Buerger’s disease and (rarely) syphilitic endarteritis may also give rise to intermittent claudication. Anaemia may precipitate or aggravate it. The physical examination of the affected limbs is usually grossly normal. However, careful examination will reveal an absence of pulsation at the dorsalis pedis and posterior tibial arteries. Similarly, the popliteal pulse may also be lost. The femoral artery pulsation may be absent, reduced or normal. There may, however, be an audible femoral bruit. After exercise, the foot may appear unduly pale; with rest, the returning flush of normal colour spreads gradually over its surface.

Similar symptoms, although usually bilateral, may also result from **spinal stenosis** *(neurogenic claudication)* (see BACK, PAIN IN, ‘Mechanical causes’, p. 47). Walking produces characteristic aching in the legs, sometimes with paraesthesias in the feet. Ankle jerks may be lost during exercise. The condition is usually distinguishable from arterial disease by a history of chronic back pain, usually accompanied by marked radiographic changes of spondylosis, and by the presence of normal peripheral leg pulses and good skin perfusion. The pathophysiology is becoming more clearly understood. Although it does not usually present until patients are elderly, narrowing of the spinal canal is a developmental defect that is associated with low birth weight, early gestation and low social class, and is also more common in the first-born child. Since the spinal canal is a ‘closed box’, any lesion that occupies the space will clearly have a greater affect in a patient who has a small-diameter canal than if the canal is normal. The condition is seen in the elderly as a result of degenerative change causing soft-tissue encroachment at multiple levels. If the compression – especially if present at two adjacent levels – is greater than that of venous pressure, venous congestion will supervene. If there are two levels of central stenosis, all the nerve roots of the cauda equina at that level will be involved. However, a single-level central stenosis, even when combined with a more distal single root canal stenosis, will only cause congestion of the single nerve root. Symptoms are probably precipitated by walking because the arterial vasodilatation in the nerve root, which normally occurs on exercise, is short-lived in these patients and rapidly fails, leading to impaired nerve root conduction. The condition must be distinguished from a narrowed exit foramen that has been further compromised by a small disc protrusion, but the latter will precipitate unilateral symptoms and signs.

In concluding this section on the causes of referred radicular pain into the legs (Box L.2) remember the spinal canal is effectively a ‘closed box’ and remembering the spinal canal is normal. The condition is due to ineffective arterial blood supply to the leg muscles, and it is caused by narrowing of the iliac, femoral or popliteal arteries. Atheroma with or without thrombosis is the usual culprit, but embolism, Buerger’s disease and (rarely) syphilitic endarteritis may also give rise to intermittent claudication. Anaemia may precipitate or aggravate it. The physical examination of the affected limbs is usually grossly normal. However, careful examination will reveal an absence of pulsation at the dorsalis pedis and posterior tibial arteries. Similarly, the popliteal pulse may also be lost. The femoral artery pulsation may be absent, reduced or normal. There may, however, be an audible femoral bruit. After exercise, the foot may appear unduly pale; with rest, the returning flush of normal colour spreads gradually over its surface.

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**Box L.2 Causes of radicular pain radiating into the lower limbs**

<table>
<thead>
<tr>
<th>Intrathecal</th>
<th>Extraspinal</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Irritation of the meninges by haemorrhage</td>
<td>• Spinal abscess – tuberculosis, brucellosis/osteomyelitis</td>
</tr>
<tr>
<td>• Infection</td>
<td>• Vertebral tumour – primary, secondary, myeloma</td>
</tr>
<tr>
<td>• Intrathecal injections</td>
<td>• Spondylolisthesis</td>
</tr>
<tr>
<td>• Hydatid cyst</td>
<td>• Vascular malformations</td>
</tr>
<tr>
<td>• Post-herpetic neuralgia</td>
<td>• Extraspinal</td>
</tr>
<tr>
<td>• Neurofibroma or other tumour</td>
<td>• Penetrating injuries</td>
</tr>
</tbody>
</table>

**Box L.3 Causes of pain in the buttock**

- Traumatic – pelvic fracture (especially due to osteoporosis in the elderly)
- Inflammatory – sacroiliitis due to spondyloarthopathies, polymyalgia rheumatica
- Infection – tuberculous or pyogenic
- Tumours – primary and metastatic tumours affecting the pelvic bones (see ‘Pain in and around the knee’, p. 382)
- Metabolic – osteomalacia, Paget’s disease of bone
- Bursitis – ischial, greater trochanteric

N.B. Referred pain from the lower lumbar spine is by far the most common cause of buttock pain, and is often associated with tenderness in the region of the posterior superior iliac spine.
noticed immediately after the injury. Postpartum pelvic pain can affect the pubic symphysis, the sacroiliac joint or both these regions of the pelvic ring. Symptoms usually settle over a few weeks, but occasionally pain – often accompanied by a feeling of instability – is persistent. Inflammatory aseptic sacroilitis may occur as part of ankylosing spondylitis, when it may be the presenting symptom, or in association with psoriasis, inflammatory bowel disease, acute anterior uveitis or Reiter’s syndrome. Sepsis of the sacroiliac joint may arise as a result of tuberculosis or pyogenic infection.

Clinical signs of sacroiliac joint disease are notoriously unreliable. Induction of local pain by forced abduction of the flexed hips or direct pressure over the sacrum may be helpful in acute cases. Other clinical tests are of little value. X-ray changes generally appear late. In patients with sepsis, other abnormalities, such as pulmonary tuberculosis or a cold abscess pointing over the sacroiliac joint or in a gluteal fold, may be found. In the acute phase, the investigations of choice are radioisotope bone scanning and CT or MRI scanning.

Osteomalacia should be suspected in malnourished individuals, and this gives rise to a characteristic stiff gait. In the elderly, Paget’s disease of bone may produce a local ache, although symptoms are often due to secondary osteoarthritis of the hip, while fractures through the softened bone can occur with minimal trauma. Deformity at the hip secondary to softening of the bone due to Paget’s disease is known as protrusio acetabuli. Ischial bursitis is a particularly troublesome condition in paraplegics who are not only wheelchair-bound, but the ischial tuberosity is very prominent due to wasting of the gluteal muscles. Skin breakdown can occur, and surgical resection of the tuberosity may be required. Greater trochanteric bursitis is common, and often misdiagnosed. This is surprising since it exhibits the classical triad of any tendinitis with: (i) local tenderness; (ii) pain on resisted active movement – in this case, resisted abduction of the hip; and (iii) pain on maximum passive movement (i.e. full hip adduction). It may be associated with underlying hip pathology and chronic mal-use of the abductors. It is also frequently secondary to other mechanical problems of the leg causing minor abnormalities of gait, for example malfunction in the patellofemoral joint in adolescents, especially females, who then compensate by overuse of the iliobibial band.

PAIN IN THE GROIN AND FRONT OF THE THIGH

Hip joint disease

Hip pain is classically felt anteriorly in the groin and frequently radiates down the front of the thigh to the knee. Occasionally, hip disease presents purely with pain in the knee. This is particularly so in the teenage disease of slipped capital femoral epiphysis. Pain in the buttock usually arises from the lumbar spine or sacroiliac joint, and is an uncommon presentation of hip pathology.

Hip joint disease in children and adolescents

Developmental dysplasia of the hip (formerly known as congenital dislocation of the hip) should be diagnosed at birth, when not only is treatment straightforward in the majority of cases but the prognosis is also good. Consequently, all practitioners with responsibility for children – doctors, nurses and health visitors – should be able to make this diagnosis. As the condition is not always evident post partum, it is good practice to examine the baby’s hip more than once, and special care needs to be taken if there is a family history or if there has been a breech delivery. There is a high incidence in communities in which the young are swaddled whereby the hips are kept adducted, and a low incidence in communities in which babies are carried on their mothers’ backs with the hips abducted. The reason is obvious since the initial treatment requires the hip to be held in flexion and abduction. In unilateral cases, signs include skin crease asymmetry and restricted abduction in flexion (Fig. L.7); Ortolani’s test demonstrates the click-clunk of reduction, while Barlow’s test demonstrates the reverse (i.e. a click as the hip is dislocated). Any ‘clicking hips’ should be examined with ultrasound, which is the investigation of choice; X-rays are much harder to interpret as the ossific nucleus of the head is not present at birth but appears during the first year. From 12 weeks, an asymmetrical appearance of the ossification centre warrants careful re-examination and an ultrasound. Bilateral dislocated hips are more difficult to diagnose as there is no asymmetry and both hips have similar restriction of movement. As a
consequence, there is a higher incidence of bilateral dislocation presenting late, often when the child starts to walk and the characteristic waddling gait becomes evident. An additional feature of bilateral dislocation is a markedly increased lumbar lordosis.

Two other conditions (both of which are rare) can produce similar physical signs in the young child. The first is infantile coxa vara, in which there is a stress fracture through the neck of femur, and as a consequence the leg is short and the hip lacks abduction due to impingement of the greater trochanter on the ilium (Fig. L.8). The head of the femur is, of course, located in the socket. The other condition results from the destruction of the head, consequent upon infection, and is termed acute epiphysitis of infancy (Tom Smith’s disease) (Fig. L.9). Close inspection will usually reveal a small scar often hidden in the groin, the hallmark of an old sinus, where the abscess plus the destroyed head of the femur was discharged.

Legg–Calvé–Perthe’s disease is usually just referred to by the name of the third describer of the condition, Perthe. There is still much dispute concerning the precipitating cause. It is probably multifactorial, but the common final pathway causes avascular necrosis of the ossific nucleus (Fig. L.10) and consequent deformation of the hip (Fig. L.11). Severe cases are likely to lead to development of osteoarthritis in young adult life (Fig. L.12). Presenting features are a painful limp and restricted hip movement, especially abduction.

Perthe’s disease accounts for only a small percentage of children presenting with an irritable

![Figure L.8](image1.png) Conditions with similar signs to a congenital dislocation of the hip. Infantile coxa vara. In the left hip, the epiphyseal plate is vertical, a triangular bony fragment is present inferiorly in the neck, and the trochanteric ossification centre is lying proximal (i.e. the head is ‘slipping off the neck’ and the leg is short).

![Figure L.9](image2.png) Conditions with similar signs to congenital dislocation of the hip. Septic arthritis: the end result is loss of the femoral head, and consequently the hip dislocates.

![Figure L.10](image3.png) Legg–Calvé–Perthe’s disease. Anteroposterior X-ray of the pelvis in a 10-year-old boy who presented with avascular change affecting the whole of the right capital epiphysis. Note the widened joint space, the flattened irregular epiphyseal plate and the widened metaphysis of the neck.

![Figure L.11](image4.png) Same patient as Fig. L.10. Five years later, the hip has healed, but with marked deformity.
hip (transient synovitis). This affects a similar age group, usually aged 6–12 years, but the symptoms settle rapidly with rest, and the hip reverts to normal. Ultrasound shows an effusion within the hip only in a minority of cases. Its importance is merely to distinguish it from other serious diseases such as infection, juvenile rheumatoid arthritis and Perthe's disease.

Slipped capital femoral epiphysis occurs around the time of puberty and therefore presents a little later in boys than in girls. In essence, it is a stress fracture occurring through the growth plate at the level of the hypertrophic cartilage cells. Many of the children are obese (Fig. L.13) or else tall and thin, and it has in the past been postulated that there could be a hormonal imbalance. However, hormone studies are normal, unless it occurs outside the age group 9–14 years in girls and 11–16 years in boys, when it may be secondary to a pituitary tumour. A slipped epiphysis may present purely with knee pain, and it is easy to confuse with the common teenage anterior knee pain secondary to problems in the patellofemoral joint, so never attribute knee pain to the knee in this age group unless you have demonstrated that hip movements are normal. In the early stages, both the symptoms and signs may be subtle. The pain and the limp may only be present on activity, and the external rotation deformity of the leg may be very mild, as are the loss of hip movements, especially flexion, abduction and internal rotation. Look in particular for increasing external rotation as the hip is flexed. The X-ray changes are easy to miss, especially on the anteroposterior film (Figs L.14 and L.15). As the degree of the slip increases, so the signs – including shortening of the leg – become more obvious. If the diagnosis is delayed, these children are particularly susceptible to a minor injury causing an acute-on-chronic slip in which the deformity at the level of epiphyseal plate suddenly increases; indeed, the head and neck can separate as if there had been a fracture. This group is the most difficult to treat and has the highest incidence of avascular necrosis.
Septic arthritis of the hip and osteomyelitis of the proximal femoral neck are both important diseases to diagnose and treat before the hip joint is destroyed. Since the capsule of the hip has ‘migrated’ down the femoral neck, the proximal metaphysis lies within the hip joint. Consequently, pus forming from infection in this region can discharge into the hip, causing secondary septic arthritis. In the classical scenario, a child presents with a high fever and is obviously ill. In modern practice, however, the disease may be less fulminating but, while it may be less lethal, it can still be sufficient to destroy the hip. The error, which the author has seen on a number of occasions, is to mistake this scenario for juvenile rheumatoid arthritis of the hip. To prevent this error, the rule is ‘treat the patient for septic arthritis and investigate to determine if it is a juvenile rheumatoid arthritis’. The child will present with pain and limp, or, if the infection is severe, will not move or use the leg. There is a fever, and hip movements are grossly restricted; indeed, any attempt at moving the hip may be very painful. X-rays will be normal since any destruction of bone will not show for 10 days. Take blood cultures in addition to the routine Hb, WBC and differential, ESR and CRP. Ultrasound will confirm an effusion which, if present, should be aspirated under ultrasound guidance and sent for culture. Remember that a ‘clear’ aspirate does not exclude infection since acute metaphyseal osteomyelitis causes a sympathetic effusion in the initial stages. Since ‘pus under pressure’ can have devastating complications – osteonecrosis, dislocation, growth disturbance, septicaemia – it is safest to proceed to open arthrotomy and lavage if the clinical symptoms, signs and ultrasound indicate infection as surgery provides ‘decompression’ as well as pus for diagnosis. With proximal metaphyseal osteomyelitis, use X-ray image intensifier control to avoid damage to the growth plate when drilling up the femoral neck.

Tuberculosis of the hip can be deceptive, and there is often a delay in making the diagnosis, particularly in Western society, now that this disease is uncommon. There is usually weight loss and often night sweats. Hip movements will be restricted in all directions and, characteristically, muscle wasting is often very marked. X-ray changes show a wide variety of patterns, primarily dependent on the initial site of the infection – that is, whether this was a tubercular synovitis, in which case there may be erosions at the capsular attachment to the bones or a tubercular osteomyelitis.

Table L.4 summarizes the causes of hip joint disease in children and adolescents in comparison to adults.

Table L.4 Causes of hip joint disease

<table>
<thead>
<tr>
<th>In children and adolescents</th>
<th>In adults</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age (years)</strong></td>
<td><strong>Disease</strong></td>
</tr>
<tr>
<td>0–5</td>
<td>Developmental hip dysplasia (congenital dislocation)</td>
</tr>
<tr>
<td></td>
<td>Infantile coxa vara</td>
</tr>
<tr>
<td></td>
<td>Acute infective epiphysitis</td>
</tr>
<tr>
<td>5–10</td>
<td>Legg–Calvé–Perthe’s disease</td>
</tr>
<tr>
<td>10–15</td>
<td>Slipped capital femoral epiphysis</td>
</tr>
<tr>
<td>At any age</td>
<td>Infection</td>
</tr>
<tr>
<td></td>
<td>• Acute osteomyelitis of the proximal femur</td>
</tr>
<tr>
<td></td>
<td>• Septic arthritis of the hip</td>
</tr>
<tr>
<td></td>
<td>• Tuberculosis</td>
</tr>
<tr>
<td></td>
<td>Inflammatory arthropathy</td>
</tr>
<tr>
<td></td>
<td>• Juvenile rheumatoid arthritis (Still’s disease)</td>
</tr>
<tr>
<td></td>
<td>Degenerative arthropathy</td>
</tr>
<tr>
<td></td>
<td>• Osteoarthritis</td>
</tr>
<tr>
<td></td>
<td>• Osteonecrosis (avascular necrosis)</td>
</tr>
</tbody>
</table>

(Continued)
Hip joint disease in adults

Trauma

Subcapital and intertrochanteric fractures of the femoral neck are not only common, but the age-related incidence also continues to rise. The underlying problem is bone fragility due to osteoporosis or osteomalacia, the latter being all too easy to overlook. They often occur after minimal trauma such as a fall or stumble. When the fracture is displaced the signs are obvious, as the leg is shortened and externally rotated, and the patient is unable to walk. Undisplaced fractures can, however, be deceptive, and if missed are often a cause of litigation (Figs L.16 and L.17). If the X-ray is inconclusive, obtain a MRI or technetium bone scan before discharging an elderly patient suffering groin pain after a fall.

Infection

Pyogenic hip infection in an otherwise fit adult is uncommon, but it is a ‘silent area’ for sepsis in intensive care units as it can develop secondary to a femoral artery or a femoral vein puncture (i.e. the needle missed the vessel and punctured the hip, which lies directly behind the vessels). Occasionally, pyogenic infection complicates an osteoarthritic joint. Adult tubercular infection is again on the increase. Surprisingly, in the adult, it can be mistaken for osteoarthritis or rheumatoid arthritis (Fig. L.18). A proper history will reveal the difference. There is

Table L.4 (Continued) Causes of hip joint disease

<table>
<thead>
<tr>
<th>Other causes</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Neurogenic</strong></td>
</tr>
<tr>
<td>• Lumbar root irritation</td>
</tr>
<tr>
<td>• Obturator nerve irritation</td>
</tr>
<tr>
<td>• Meralgia paraesthetica</td>
</tr>
<tr>
<td><strong>Abdominal pathology</strong></td>
</tr>
<tr>
<td>• Hernias: inguinal, femoral, obturator</td>
</tr>
<tr>
<td>• Retroperitoneal inflammation</td>
</tr>
<tr>
<td><strong>Infective</strong></td>
</tr>
<tr>
<td>• Local abscess</td>
</tr>
<tr>
<td>• Infiamed inguinal lymph nodes</td>
</tr>
<tr>
<td>• Psoas abscess</td>
</tr>
<tr>
<td><strong>Tumours</strong></td>
</tr>
<tr>
<td>• Metastatic disease involving the pelvis or proximal femur</td>
</tr>
<tr>
<td>• Primary malignant bone tumours: osteosarcoma, chondrosarcoma</td>
</tr>
<tr>
<td>• Benign bone tumour and tumour-like conditions</td>
</tr>
<tr>
<td>• Soft-tissue tumours: pigmented villonodular synovitis</td>
</tr>
<tr>
<td><strong>Muscle and tendon</strong></td>
</tr>
<tr>
<td>• Sprains (e.g. adductor tendon sprain)</td>
</tr>
<tr>
<td>• Haematoma: myositis ossificans</td>
</tr>
<tr>
<td><strong>Other pathology</strong></td>
</tr>
<tr>
<td>• Synovial chondromatosis</td>
</tr>
</tbody>
</table>

Figure L.16 Subcapital fracture of the femoral neck. This 70-year-old woman presented with mild groin pain and a limp following a fall. The anteroposterior X-ray was erroneously considered normal, although there is a suggestion of a fracture in the calcar (arrows).

Figure L.17 Same patient as Fig. L.16. Shortly after discharge, the patient stumbled, and the fracture was displaced.

Figure L.18 Tubercular synovitis of left hip. This was initially diagnosed as a monoarticular variant of rheumatoid arthritis. In addition to joint space narrowing, there is a marked bony erosion at the synovial margins, most marked in the acetabular fossa (arrow).
often marked gluteal or quadriceps wasting, as well as signs of hip irritability, weight loss and night sweats (see ‘Hip joint disease in children and adolescents’, above). Radiologically, tubercular synovitis can be confused with pigmented villonodular synovitis, which also causes bone erosion at the synovial margins (see ‘Pain in and around the knee’, p. 362, Fig. L.36). Chronic infection, often with sinus formation around a total hip replacement, is a ‘new’ disease of modern medicine. It is important to establish a precise bacteriological diagnosis as a wide variety of organisms has been implicated. The infection rate for primary hip arthroplasty should be one per cent or less.

Inflammatory arthropathies
Although rheumatoid arthritis predominantly affects the small peripheral joints (e.g. the hands, wrists, feet and ankles), the hip is by no means spared, but the diagnosis is usually obvious due to the generality of the disease. Isolated hip joint involvement does occur with other inflammatory arthropathies, especially ankylosing spondylitis (see also p. 50). The fixed flexion deformity of the hip is a major contributory factor to these patients’ inability to stand erect and to see ahead (see Figs B.3 and B.4).

Degenerative arthropathy
In osteoarthritis, typically the symptoms of pain and stiffness are slowly progressive, usually over many years. The hip joint in particular has a propensity to cause night pain, which is characteristically felt in the groin with radiation down the front of the thigh. While movements of the joint are restricted in all directions, close inspection will reveal that this is not uniform. Abduction and internal rotation are usually more restricted than adduction and external rotation, although sometimes this pattern may be reversed. A useful clinical definition of an osteoarthritic joint is that there is a free painless arc of movement terminated by painful restriction. X-ray changes are also slowly progressive (Fig. L.19). A patient with X-ray changes that could be mistaken for osteoarthritis but clinically has excessive movement is likely to have a Charcot joint (Figs L.20 and L.21).

Beware of the joint presenting with marked muscle spasm and very painful grossly restricted movements. Even if the X-rays show signs of osteoarthritis, this is not the diagnosis until proven otherwise – septic arthritis, gout, chondrocalcinosis (pseudo-gout) and other inflammatory arthropathies require exclusion.

In avascular necrosis (osteonecrosis), the articular cartilage is normal, and the problem is in the underlying bone. The pattern of vascular supply to the head of the
femur leaves the hip particularly prone to avascularity. Interestingly, osteonecrosis always affects the convex side of the joint; other sites are the capitulum, the proximal pole of the scaphoid, the dome of the talus and the navicular. Many diseases are associated with osteonecrosis of the hip, which should therefore never be regarded as a primary diagnosis. The main groups are: trauma, especially following fractures through the femoral neck; alcoholism (Figs L.22–L.24); steroids; haemoglobinopathies (especially sickle-cell disease); Caisson disease (Fig. L.25); and iatrogenic conditions (i.e. secondary to treatment for childhood congenital dislocation of the hip, adolescent slipped capital femoral epiphysis, etc.). In the early stages, osteonecrosis can be very deceptive as the pain may be mild and atypical, presenting in the thigh or the knee, or with discomfort around the greater trochanter rather than groin pain. Movements may be largely preserved, but some restriction will be evident on close inspection. The X-ray in the early stages is often normal, and only later does the sclerosis in the femoral head become evident (this is indicative of new bone being laid down on dead trabeculae). A technetium bone scan and an MRI scan are both sensitive investigations in the early stages. The presence of a subchondral fracture through the avascular bone (revealed as the ‘crescent sign’ on the lateral plain X-ray) is a poor prognostic sign, as this means that the overlying cartilage plus a slither of the subchondral bone plate has separated from the femoral head.

Figure L.21 Same patient as Fig. L.20. Anteroposterior X-ray of the lumbar spine and sacrum.

Figure L.22 Osteonecrosis of the right hip secondary to alcoholism in a 30-year-old man. Anteroposterior X-ray showing increased sclerosis of the right femoral head.

Figure L.23 Same patient as Fig. L.22. Lateral X-ray suggests sclerosis is localized to the anterior aspect of the head (shown between two arrows).

Figure L.24 Same patient as Fig. L.22. The presence of sclerosis is confirmed by a computed tomography scan.
Other conditions causing groin and anterior thigh pain

Radicular pain arising from the nerve roots of the lumbar plexus is much less common than with the sacral plexus, since degenerative back disease is far more frequent at the L4/L5 and L5/S1 levels than the upper lumbar spine. Indeed, beware of attributing pain to an upper lumbar disc since this is relatively uncommon and there is, therefore, a greater chance of the patient having a more serious entity such as a benign spinal tumour or irritation of the plexus by retroperitoneal pathology with pain radiating down the femoral nerve anteriorly into the thigh. The signs are often subtle. Careful examination may find wasting of the quadriceps. The knee jerk is rarely lost as it has a dual innervation from L3 and L4 (see Table L.3, p. 351).

Obturator nerve irritation produces pain radiating along the inner side of the thigh. It is uncommon, and tends to be associated with serious disease – metastatic disease within the pelvic cavity or within the pelvic bones, especially the pubis. Physical signs are often minimal, but there may be wasting of the adductor group.

Meralgia paraesthetica is common but frequently missed. It is due to entrapment of the lateral cutaneous nerve of the thigh, where it exits through the deep fascia, just distal to the inguinal ligament below the anterior superior iliac spine and close to the origin of the sartorius. The pain is often rather diffuse, but careful testing usually reveals an area of numbness anterolaterally, and there is often a tender spot at the exit point of the nerve through the deep fascia.

Long-acting local anaesthetic injections to this site will not only confirm the diagnosis but are also often curative.

Local causes of infection in the groin, including lymphadenitis of the inguinal nodes, usually have obvious signs. Inspect the leg for a possible primary source of infection and, if present, remember to check for diabetes. The incidence of spinal tuberculosis is rising rapidly worldwide due to the HIV/AIDS epidemic, and one can expect the incidence of psoas abscess will once again increase. The spinal pathology is usually at the thoracolumbar junction, with the caseous material tracking along the psoas sheath to the groin.

By far the most common tumours causing pain around the hip and down the thigh are metastases, destroying either part of the pelvis or the proximal femur. A useful rule states that ‘if 50 per cent of the femoral cortex is involved, then there is a 50 per cent chance of fracture’. Primary malignant tumours of bone are very uncommon, but when they do occur the region of the hip joint is the second most common site after the knee (see ‘Tumours of bone’, p. 371). Chondrosarcomas in particular arise from the flat bones, the pelvis and the scapula, often from a pre-existing exostosis (see Fig. L.44, p. 373). Consequently, any change in size in an exostosis in adult life needs urgent investigation. Chondrosarcomas are slow-growing and usually affect the 30- to 50-year age group.

Pigmented villonodular synovitis, which is now classified as a benign soft-tissue neoplasm (see ‘Pain in and around the knee’, ‘Arthropathies’, p. 366), presents as a low-grade inflammatory condition. In the hip, it can mimic rheumatoid arthritis and tuberculous synovitis due to the presence of periarticular erosions on X-ray. Eventually, the repeated haemarthroses lead to severe osteoarthritis.

Muscle and tendon sprains around the hip are common, especially in athletes. Strains of the attachments of the adductors are probably the most frequent, but all muscle groups including the psoas are at risk. Small inguinal hernias are the main differential diagnosis. Myositis ossificans (Figs L.26 and L.27) results from a direct blow to the thigh and consequent ossification of a haematoma in vastus intermedius. The loss of knee flexion distinguishes it from a malignant bone tumour in which knee mobility is retained (see ‘Tumours of bone’, p. 371). The condition resolves over several months. Myositis ossificans is a well-recognized complication of hip surgery, especially incisions that damage the periosteum of the outer iliac wall. The adductors, gluteus medius and gluteus minimus are the most frequent sites. Minor degrees
resolve, but when severe, loss of hip movement is permanent without further surgery.

The hip is the third most common joint, after the knee and elbow, to be affected by synovial chondromatosis (Figs L.28 and L.29). It usually presents as a mild synovitis, with a tendency for the hip to give way. Actual locking is uncommon. CT and MRI scans will clearly delineate loose bodies, which are easy to overlook on plain X-rays.

Figure L.26 Myositis ossificans of the left quadriceps femoris in a 22-year-old man. Lateral X-ray of the left thigh. There is an area of ossification lying anteriorly over the mid shaft of the femur. Note the area of separation between the myositis ossificans and the femur.

Figure L.27 Same patient as Fig. L.26. The ossification is more clearly seen on the computed tomography scan.

Figure L.28 (a) Synovial chondromatosis of the left hip. Anteroposterior and lateral X-rays of the left hip. Numerous loose bodies are present. (b) Loose bodies are confirmed by computed tomography scanning; the majority appear to be lying anteriorly, but some are present deep within the joint.

Figure L.29 Synovial chondromatosis of the knee. Numerous small cartilaginous loose bodies approximately 3–5 mm in size, found at a diagnostic arthroscopy for chronic synovitis in a 22-year-old man. The X-ray was normal.
PAIN IN AND AROUND THE KNEE

The knee is a complex joint both as regards: (i) its anatomy – in practical terms, it is composed of three joints, the patellofemoral joint and the medial and lateral compartments of the tibiofemoral joint; (ii) its mechanics; and (iii) the presenting diseases/injuries, which together cover almost the entire range of musculoskeletal conditions. The objective in taking the history is to narrow the options. White's schema (Table L.5) includes the six most common categories of knee problem and is a helpful practical approach. There are numerous specialized physical

| Table L.5 The six categories of knee pain (after White, S., Arthritis and Rheumatism Council for Research: Practical Problems No. 10, Series 3 'The Assessment and Management of Knee Problems') |
| --- | --- | --- |
| 1 Internal derangement/‘locked’ knee | Meniscus  
- vertical tears  
- horizontal cleavage tears  
- degenerate medial meniscus syndrome | Loose bodies  
- osteochondral fractures  
- osteochondritis dissecans  
- synovial chondromatosis  
- osteoarthritis | Articular surface irregularity  
- chondral flaps  
- osteoarthritis with severe eburnation |
| 2 Ligament injuries | Medial collateral  
- complete tears  
- partial tears | Lateral collateral  
- complete tears  
- iliobial band avulsion | Anterior cruciate  
- Posterior cruciate |
| 3 Arthropathies | Osteoarthritis: varus knee (medial compartment) | Osteoarthritis: valgus knee (lateral compartment) | Inflammatory arthropathies  
- rheumatoid arthritis  
- chondrocalcinosis gout  
- Haemophilia |
| 4 Patello-femoral derangement/ extensor mechanism injuries | Anterior knee pain syndrome  
Chondromalacia patellae  
Lateral hyperpressure syndrome | Patellar instability  
- subluxation  
- dislocation  
- acute  
- recurrent  
- habitual  
- permanent | Extensor mechanism injuries/stress conditions  
- Osgood–Schlatter’s disease  
- ruptured patellar tendon  
- Johanssen–Larsen syndrome/  
- patellar fractures  
- quadriceps femoris injuries  
- myositis ossificans |
| 5 Infection/tumours/tumour-like conditions | Pyogenic infection  
- septic arthritis  
- osteomyelitis | Chronic infection  
- tuberculosis  
- infected arthroplasty  
- Brodie’s abscess | Tumours  
A. Malignant  
- osteosarcoma  
- Ewing’s sarcoma  
- chondrosarcoma  
B. Benign  
- osteoelastoma  
- osteochondroma  
- chondromablastoma  
- enchondroma  
- chondromyxoid fibroma |
| 6 Bursitis/cysts/tendinitis | Bursitis  
- prepatellar  
- infrapatellar, etc. | Cysts  
- semimembranosus  
- Baker’s cyst  
- meniscal | Tendinitis  
- iliobial band insertion  
- pes anserinus insertion  
- semimembranosus insertion  
- biceps femoris insertion |
signs, and a good history will enable one to select the correct set. All patients should be examined standing to observe any malalignment (Figs L.30 and L.31), walking to observe the gait pattern, and then lying down on a firm couch (examining the knee on a soft bed makes it much more difficult to determine whether there is loss of extension; this is an important physical sign as full extension is a requisite for normal knee function).

Two warnings! First, the knee and the hip are supplied by the same nerves, and hence hip pathology can present purely with knee pain. The classic condition in this respect is slipped capital femoral epiphysis of the teenager. It is all too easy to dismiss this as teenage anterior knee pain. Second, ensure that any pathology found on special investigations such as arthroscopy and MRI scans actually matches the patient's symptoms. Degenerative change affecting the menisci and the patellofemoral joint is almost universal over the age of 40 years, and the consequent changes on scans usually lie within the spectrum of normality.

**Internal derangement (locked/locking knee)**

The characteristic of these knees is that they suddenly jam, and the joint cannot fully extend. The most common cause is meniscal tears, followed by loose bodies, while chondral flaps are relatively uncommon and have only been appreciated since the advent of arthroscopy. Transient episodes of ‘catching’, in which the fragment catches and then releases, often present as ‘giving way’. A locked knee needs to be distinguished from other acute causes of loss of extension such as a haemarthrosis or acute arthritis, especially gout or pseudo-gout (chondrocalcinosis). A more difficult differential is from a patellar dislocation. There are two main types of meniscal tear: vertical tears, which occur more frequently in the medial than the lateral meniscus, are usually secondary to a twisting injury sustained while playing sports. As with other knee injuries, they are much more common in men than women, although there is a rising incidence in the latter now that girls have started to play football and rugby. Vertical tears...
usually begin on the inferior aspect of the meniscus in the posterior horn. With recurrent injuries, the tear spreads, until the loose fragment – the ‘bucket handle’ – eventually becomes sufficiently mobile to displace into the intercondylar notch and hence lock the knee. In keeping with the pathology, the classical history is one of recurrent episodes of pain affecting the medial side of the knee precipitated by a twist and usually accompanied by a transient effusion. Symptoms settle over a few days, and sports can then be resumed. As the tear enlarges, so less trauma is required to produce symptoms, and patients often complain that the joint feels insecure. Eventually, giving way on minor provocation occurs, followed finally by true locking. Examination between episodes may fail to reveal any abnormality. When symptomatic, there is usually a small effusion, tenderness is present along the appropriate joint line, and tibial rotation is painful. The classical special manoeuvre is the McMurray test (tibial rotation with the joint under compression), in which the examiner endeavours to reproduce not just the pain, but also a ‘clunk’ as the meniscal fragment dislocates. Check, in particular, the integrity of the ligaments as the meniscus is at much greater risk of being torn if the knee is unstable.

Horizontal cleavage tears occur in older age groups, generally 30–60 years. By comparison with the vertical tears, the initial trauma is less severe; indeed, the patient may have no recollection of any significant trauma. Otherwise, the history is similar with recurrent episodes, usually precipitated by a minor twist. The tear gradually increases, and eventually a ‘parrot beak flap’ can dislocate between the condyles, causing locking or giving way.

A very different presentation of the wearing meniscus is the immobile or degenerative medial meniscus syndrome. This is also usually precipitated by a minor twisting injury in late middle age (50–65 years), but symptoms are often persistent, becoming aggravated by the slightest twist, for example going around a corner. If severe, it may be accompanied by giving way, but there is no mechanical history of locking. The medial joint line is markedly tender and acute pain can be precipitated by tibial rotation. Characteristically, patients are often woken at night and complain that they cannot sleep on their side as they are unable to rest one knee on top of the other. In most instances, there is a horizontal cleavage tear of the medial meniscus. This syndrome usually responds well to conservative treatment. The features reflect the sensory supply to the meniscus, which is predominantly at its attachment to the capsule. As the meniscus loses its natural pliability, so there is increased tension exerted peripherally.

The knee is the most common site for loose bodies. These are frequently seen on plain X-rays, but only a minority are intra-articular and therefore a cause of mechanical symptoms. Debris within a joint usually becomes attached to the synovium, the cells of which then proliferate around the loose body, not just entrapping it but extruding it into the subsynovial layer. Even when loose bodies are theoretically intra-articular, they may be ensnared in a recess and hence are unable to float free within the joint cavity; an example is loose bodies in a Baker’s cyst. There is therefore a tendency to overdiagnose loose bodies as a cause of symptoms. They precipitate locking or giving way, but are not a cause of persistent pain. Osteochondritis dissecans affects adolescents, and it is always on the convex surface of a joint. For example, in the knee, it affects a femoral condyle, in the elbow the capitulum, and in the ankle the dome of the talus. It is best regarded as a shear fracture through the subchondral plate. The important feature is that the overlying articular cartilage is normal and, as the osteochondral fragment does not separate to become...
a loose body until late in the course of the disease, the diagnosis is easily missed at arthroscopy since the articular surface is often still intact. Prior to separation, complaints are often of rather vague symptoms, including a feeling of insecurity. There is usually a slight but definite loss of full extension; this is a particularly helpful sign in distinguishing osteochondritis dissecans from teenage anterior knee pain. The lesion shows clearly on a technetium bone scan and MRI (Fig. L.33), but it is easy to overlook on plain X-rays.

Synovial chondromatosis is a rare condition in which a change in the synovium leads to the production of multiple loose bodies. There are two forms, the osteo-chondral and the cartilaginous ‘rice body’ types. In the former, there are multiple discrete loose bodies, usually numbering between 10 and 20, and approximately the size of peas. They are visible on plain X-rays, and cause locking. In the second type, there are hundreds of small loose bodies, which, being cartilaginous, are invisible on X-rays (see Fig. L.29). These present with persistent aching discomfort, accompanied by an effusion and synovial thickening. They do not cause locking. Clinically, they are indistinguishable from other low-grade forms of chronic synovitis. These loose bodies usually appear as a surprise on arthroscopy.

Ligament injuries

Everything about the ligaments of the knee is complex – their anatomy, the mechanism of their injuries, the precise diagnosis of those injuries, and their treatment.

Note:

- The more serious the injury, the less pain.
- There is a high rate of popliteal artery damage associated with posterior dislocations of the knee. Rule: ‘if a posterior dislocation is suspected, an angiogram is mandatory.’

Contrary to what one might expect, these are not all high-velocity injuries; indeed, the author can clearly recall a case in which an overweight woman aged 70 ruptured both cruciate ligaments and transected her popliteal artery merely by tripping over a small step. In the classical triad – injury with rupture of the medial ligament, rupture of the anterior cruciate ligament and a tear of the medial meniscus – the knee has been subject to a severe valgus external rotatory force, for example a rugby tackle. At the moment of injury, there is a severe pain often accompanied by a loud crack such that the individual may fear the leg has ‘broken’. After a few minutes, the pain has largely disappeared, and many players then seek to carry on playing. However, the next time they receive a ball, the knee gives way as soon as they pivot or twist. When first seen, the injury does not appear serious, as there is little or no pain (the ligaments having snapped, the nerve endings do not fire since they respond to tension) and there is no haemarthrosis (the capsule has been disrupted and the blood has escaped into the soft tissues). Finally, if the athlete is fit, their muscles are strong, and hence the doctor may not detect the instability.

First, examine the uninjured knee and not only determine the degree of laxity in both full extension and in 30° of flexion, but in addition note the feel of the ‘end point’ while at the same time palpating the ligament; this will appear as a well-defined taut band. Repeat this on the injured knee. If, on the medial side, an end point cannot be felt and a taut band cannot be demonstrated,
As tibial rotation at the extreme of extension tightens locking. Anatomically, the screw-home mechanism of the knee lacks full extension. This, however, is 'pseudo-of the medial meniscus, since the pain is felt medially and confused with a locked knee due to a bucket-handle tear missed because they are very painful, but they are easily very differently from complete ruptures. These are never onset of swelling due to a haemarthrosis.

or felt a ‘pop’ from within the joint, and there is a rapid knee. Two other characteristics of the acute anterior ways of sustaining the injury – when the ski bindings the femur with a ‘jerk’ at approximately 30° of flexion. When the result is positive, the tibia will relocate onto the lateral malleolus; exert a proximal force up the leg to replicate the ground reaction force; place the left hand behind the fibula and apply an internal rotation change direction. To perform the test on the right knee, run in a straight line, but the knee collapses when they change direction. To perform the test on the right knee, lift up the extended leg by the heel with the thumb over the lateral malleolus; exert a proximal force up the leg to replicate the ground reaction force; place the left hand behind the fibula and apply an internal rotation force; still applying these forces, gently flex the knee. When the result is positive, the tibia will relocate onto the femur with a ‘jerk’ at approximately 30° of flexion. This test therefore reproduces one of the common ways of sustaining the injury – when the ski bindings fail to release and the ski applies a twisting force to the knee. Two other characteristics of the acute anterior cruciate rupture are that the patient may have heard or felt a ‘pop’ from within the joint, and there is a rapid onset of swelling due to a haemarthrosis.

Partial tears (sprains) of the medial ligament present very differently from complete ruptures. These are never missed because they are very painful, but they are easily confused with a locked knee due to a bucket-handle tear of the medial meniscus, since the pain is felt medially and the knee lacks full extension. This, however, is ‘pseudo-locking’. Anatomically, the screw-home mechanism of the knee enables one to stand without using muscle power except for slight activity in the tensor fascia lata, as tibial rotation at the extreme of extension tightens the ligaments. This mechanism is blocked by the pain from stretching the damaged medial ligament, and hence the knee lacks full extension and appears locked. The distinguishing physical sign between these two injuries is that the tenderness from a medial ligament sprain is always localized above the joint line, whereas, with a bucket-handle tear, the tenderness is along the joint line. Medial ligament sprains may take several weeks to resolve. This is particularly the case if an area of calcification forms at the femoral attachment.

MRI is the investigation of choice in ligament injuries and often provides remarkably accurate visualisation of the tear (Fig. L.34b). However, a word of warning! MRI can both overstate and occasionally obscure injuries around the knee. By itself a MRI diagnosis by no means provides an automatic indication for surgical treatment. This is especially the case with horizontal meniscal tears, acute cruciate injuries, osteoarthritis and patellofemoral problems. In particular, beware of the ‘MRI bone bruise’ which can make minor injuries seem major on obscure fractures of clinical importance.

Arthropathies

Osteoarthritis of the knee is an extremely common condition throughout the world. It can affect the medial compartment in association with a varus deformity, the lateral compartment in association with the valgus deformity, or the patellofemoral joint in association with patellofemoral maltracking. The history of medial or lateral compartment osteoarthritis is of slow deterioration over many years. Pain is brought on by activity and is usually localized to the affected side (for patellofemoral osteoarthritis, see ‘Anterior knee pain/patellofemoral problems’, below). Night pain, unlike in the hip, is not a particular feature. Acute episodes of pain, often associated with episodes of catching or giving way followed by an effusion, can occur in advanced disease. This is due to gross eburnation and irregular ridging in the affected compartment – in effect, the femur has jumped a groove on the tibia.

On examination, first check for deformity with the patient standing (i.e. bow leg, knock knee, loss of full extension; bow legs are easy to overlook in the obese). Observe the patient walking – in advanced medial compartment disease, the knee collapses into more varus. Joint line tenderness is often slight, but usually localizes to the affected side. The presence of an effusion is very variable but is often small. Movements are restricted, but this is more marked in Caucasians than in Middle Eastern or Asian populations.

The confirmatory sign for osteoarthritis is to elicit bony crepitus from the affected compartment.
For a varus knee, first stress the lateral compartment applying a valgus stress. Compress the surfaces together while flexing and extending the knee. This surface will be relatively intact, and the feeling elicited is normal smooth movement. Repeat this with a varus stress. Flex the knee through the complete range of flexion/extension, and if there is eburnation at some point through this arc, often between 40° and 60°, one not only elicits bony crepitus, but also reproduces the patient’s pain. Routine X-rays are misleading, particularly taken with a patient lying down. A standing anteroposterior X-ray usually tells the truth, either a narrowing or absence of the joint space. Even more useful is the anteroposterior film taken in 30° of flexion (Fig. L.35). In keeping with the physical signs of the ‘agony test’ described above, this can reveal total obliteration of the joint space when the standing film in extension shows only a partial loss. In summary, there is a tendency to underestimate the severity of this common condition as night pain is unusual, and the signs may be subtle and difficult to elicit, while X-rays often fail to show the true extent of cartilage loss.

The knee joint is affected in many of the inflammatory arthropathies; these are considered in detail under JOINTS, AFFECTIONS OF (p. 319). Rheumatoid arthritis characteristically results in a valgus deformity with concomitant external rotation of the tibia. Chondrocalcinosis occurs more frequently in the knee than in any other joint. It is due to the deposition of calcium pyrophosphate crystals within the menisci or within the articular surfaces. It is a very frequent X-ray finding and, in the majority of instances, is asymptomatic. Symptoms are precipitated when, secondary to early wear, pyrophosphate crystals ‘escape’ from the articular surface or from the menisci and cause an acute painful reaction within the joint. Characteristically, the knee is warm with a marked effusion and restricted mobility. Often there is a loss of range of movement.

For a varus knee, first stress the lateral compartment applying a valgus stress. Compress the surfaces together while flexing and extending the knee. This surface will be relatively intact, and the feeling elicited is normal smooth movement. Repeat this with a
of 20–30° of extension, with flexion to only 90°. Attacks of gout affecting the knee, albeit uncommon, are all too easy to misdiagnose. Confirmation of both chondrocalcinosis and gout is by finding the crystals in a synovial aspirate.

Although the diagnosis of haemophilic arthropathy is usually clear from the history, it can be difficult if it is the presenting symptom especially when the extent of the bleeding is mild. The appearances are then of a persistent synovitis with recurrent flares. Any joint can be involved, but the knees, ankles and elbows are particularly at risk. A clotting screen will elucidate the precise defect. When this is normal, a MRI scan can be very helpful in diagnosing a local cause, for example a haemangioma of the synovium or pigmented villonodular synovitis. A haemarthrosis as a complication of osteoarthritis used to be rare, but is becoming more frequent due to the increase in prescribing low-dose aspirin (see JOINTS, AFFECTIONS OF, p. 319).

Pigmented villonodular synovitis is now classified as a benign soft-tissue neoplasm. It is included in this section because it presents as a low-grade synovitis with episodic acute flares due to bleeding into the joint. It therefore mimics gout, chondrocalcinosis, monoarticular rheumatoid, haemophilic synovitis and bleeding from an arteriovenous haemangioma involving the synovium. The common sites are the knee and the hip. The knee X-rays are usually normal, but in the hip periarticular bone erosion can mimic tuberculosis. The appearances on both MRI and arthroscopy are usually diagnostic (Fig. L.36). An identical histological lesion, giant-cell tumour of the tendon sheath, occurs in the flexor tendon sheaths of the fingers. This presents as a soft-tissue swelling, or occasionally as a trigger finger (see ‘Conditions affecting the hands and fingers’, p. 717).

PAIN IN THE FRONT OF THE KNEE

Anterior knee pain/patellofemoral problems

The common cause of pain in the front of the knee is malfunction in the patellofemoral joint. It is important first to ensure that the pain is not referred from the hip, and second, that the patellofemoral joint is not overlooked as the source of pain as it can mimic several other conditions. Classically, pain from this joint radiates down anteriorly over the patellar tendon and the anterior proximal tibia. Frequently, it spreads anteromedially, and when this is combined with episodes of giving way due to patella subluxation, it mimics a torn medial meniscus. The patellofemoral joint can reproduce locking, giving way, instability, feelings of insecurity and pain from almost any other mechanical problem in the knee. Furthermore, malfunction of the extensor mechanism is also a potent cause of chronic tendon sprains in the lower limb as other muscle groups overcompensate; for example, overactivity in the iliobibial band/gluteal muscles may precipitate pain at the tibial insertion or the greater trochanter. Other sites of secondary tendinitis are the pes anserine insertion (sartorius, gracilis, semitendinosus), semimembranosus at the posteromedial corner and the biceps femoris over the fibula. With all of these tendinitides, look closely at the quadriceps/patellofemoral mechanism for a possible primary cause. Classically, patellofemoral symptoms are aggravated by taking weight through the flexed knee (e.g. on stairs, or by squatting). Indeed, if the patient can squat and ‘duck walk’ (walking in the full squat position), this mechanism is probably innocent. The exception is when the patella dislocates on extension since the patella stabilizes in the groove on flexion. Fortunately, dislocation in extension is one of the more obvious patellofemoral entities to diagnose, as the patella literally jumps laterally out of the femoral groove as the knee nears full extension.

On examination, look at limb alignment and patellar tracking. Tenderness is usually located along the medial border of the patella and over the anteromedial aspect of the knee in the triangle bounded by the medial femoral condyle, the medial tibial plateau and the patellar tendon. Test for patellar laxity: with the knee extended, feel particularly for the end point on lateral stressing. Feel for patello-femoral crepitus by placing a hand over the patella as the patient crouches. Elicit whether the patellofemoral pressure test reproduces...
the patient’s symptoms (with the knee extended, press above the proximal pole of the patella with two fingers and then get the patient to contract the quadriceps while maintaining this downward pressure).

As indicated in White’s schema, there are a variety of patellofemoral syndromes. These reflect the anatomical peculiarities of this joint. The patella: (i) being a sesamoid bone is dependent on muscular control, particularly the distal fibres of vastus medialis, to maintain its stability as it slides, tilts and rotates in spiralling down the femoral groove; (ii) has a complex shape; and (iii) has the thickest and softest articular surface of any joint.

The anterior knee pain syndrome predominantly affects teenagers, and is increasing in incidence particularly in Western society. There is a suspicion that it affects those who are athletic and those who take no exercise at all. It is the most common cause for discharge on medical grounds from the Armed Services. Fortunately, for most teenagers, the condition is mild and usually settles within 2 years. Many factors have been implicated, but it is probably best regarded as the temporary imbalance between the muscular control of the patella, the stress to which the joint is subjected and minor abnormalities in the complex anatomical shape of both the patella and the femoral groove, and the fact that the knee accounts for most of the growth in length of the lower limb.

To diagnose chondromalacia patellae, the articular surface of the patella must be shown to be excessively softened or disrupted (i.e. by arthroscopy or open surgery). There are important features distinguishing between chondromalacia and osteoarthritis. The former affects just one side of the joint, the convex patellar side, whereas osteoarthritis results in damage to both articular surfaces.

Chondromalacia on the medial facet is of little clinical importance; in the central area it can cause anterior knee pain, while on the lateral facet it is a precursor of osteoarthritis – the lateral hyperpressure syndrome.

Patellofemoral osteoarthritis is a very common condition. It can be isolated or be present in conjunction with either medial or lateral compartment osteoarthritis. In the majority of cases, the symptoms are either mild, or overshadowed by the tibiofemoral wear. When symptomatic, it causes anterior knee pain that is aggravated by taking weight on the bent knee, difficulty in kneeling or in rising from sitting, and giving way. On examination, the patella is often laterally placed and locally tender, and there is marked bony crepitus, which is frequently audible on crouching.

The two instability syndromes of patellar subluxation and patellar dislocation are distinguished from the other patellar symptoms by one feature in particular, episodic major giving way; in this, the patient is usually unable to catch himself and consequently falls. The episodes are often precipitated by a slight twist and, if they occur while walking down the stairs or crossing the road, can result in lethal injury. Ligament laxity is frequently present. This can be generalized, affecting all joints, or localized just to the patella. In between episodes of dislocation, the knee usually behaves normally and the physical signs are subtle. Feel in particular for the end point on displacing the patella laterally. This manoeuvre often precipitates a positive apprehension test (i.e. the patient recognizes that the patella is about to dislocate and the quadriceps suddenly tighten). Recurrent dislocation may also occur in a previously normal knee that was subject to a traumatic dislocation; that is, the patella was forcibly knocked out of its groove. Often the patella relocates immediately the knee is straightened. The patient is usually unaware of what exactly happened, and on inspection the only abnormal finding is that the knee is rather bruised and swollen. Routine anteroposterior and lateral X-rays are normal, and the patient is dismissed. However, a skyline view of the patella often reveals an avulsion fracture, indicating that vastus medialis has been torn from the patella. If this is not repaired surgically, there is a high incidence of recurrent dislocation. Finally, there are two other patterns of patella dislocation: habitual dislocation, in which the patella dislocates every time the knee is flexed, and permanent dislocation, which means precisely what it says – the patella is always dislocated. Remarkably, this diagnosis is frequently missed (Fig. L.37). The signs are that the knee is locked at 30° of flexion, the foot is externally rotated, and a skyline view demonstrates that the patella is not in its groove. Note that, with all patterns of dislocation, patients frequently say that the patella has dislocated medially because the deformity they notice is the uncovered prominence of the medial femoral condyle.

**Other conditions and injuries of the extensor mechanism (listed from distal to proximal)**

Osgood–Schlatter’s disease is due to a chronic stress at the tibial apophysis and occurs predominantly in teenage boys. The pain is localized around the tibial tubercle, which often becomes prominent (Fig. L.38). Symptoms usually settle over 6 months, but may last up to 2 years. Occasionally, symptoms persist after the end of growth due to a small fragment of the bone failing to unite on to the tibial shaft. The condition must
be distinguished from an acute traumatic avulsion of the tibial tubercle, which, if left untreated, results in a quadriceps lag and a fixed flexion deformity. Fortunately, this injury is rare.

Traumatic rupture of the patella tendon occurs in middle age. There is usually a small fragment of bone, avulsed from the distal pole of the patella. Surprisingly, it can be missed, since the quadriceps expansion may still be intact, and consequently close inspection is required to detect that the patella has moved proximally. Straight leg raising will be either impossible or there will be a marked lag.

Johannsen–Larsen syndrome is another cause of teenage anterior knee pain. This a chronic stress syndrome affecting the proximal attachment of the patellar tendon, the adult equivalent being jumper’s knee, which is usually seen in fading middle-aged athletes who take up such sports as marathon running. To localize the tenderness to the distal pole of the patella, flex the knee to 90°, stabilize the foot by sitting on the toes, and ask the patient to straighten the knee (i.e. put the quadriceps into action).

There are two main types of patella fracture. Stellate fractures occur from a direct blow to the patella, while transverse fractures are caused by the patient tripping and then the quadriceps exerting a sudden reflex contraction – consequently, the patella has snapped in two before the knee hits the ground. A similar mechanism causes avulsion injuries of the quadriceps or one of its component parts from the patella. This injury is often associated with chronic illness requiring steroids, renal dialysis, etc. Rupture of the rectus femoris occurs in the mid-thigh region and is an injury of athletes. When the quadriceps contracts, the rectus then rolls up like a ball. This condition must be distinguished from soft-tissue tumours such as an intrafascial lipoma. Myositis ossificans affects the vastus intermedius. There is usually a clear history of a blow to the quadriceps. The resulting haematoma then ossifies (see Figs L.26 and L.27). The diagnosis becomes obvious with the passage of time, but it can be a confusing condition in the early stages when the thigh is merely swollen and X-rays are normal. Evidence of calcification appears after 6 weeks. The differential diagnosis is from a tumour. The important distinguishing clinical sign is that, with tumours of bone, the knee remains mobile, whereas with myositis ossificans or infection, movement is lost. With myositis ossificans, the area of ossification gradually remodels, and knee movements return over a period of a year to 18 months.

Infection
The knee is the most common site for pyogenic osteomyelitis. The infection begins on the metaphyseal side of the growth plate in either the distal femur or proximal tibia. The child presents with pyrexia and severe pain, and any movement of the knee is extremely painful. There is usually a small effusion present, particularly if the infection is in the distal femur. X-rays are normal in the early stages and do not become abnormal until pus, classically due to Staphylococcus aureus, has formed, broken through the cortex and

Figure L.37 Permanent dislocation of the patella accounts for this remarkable X-ray (a) in an 18-year-old with juvenile rheumatoid arthritis. When the patella was re-sited (b), the femur and tibia realigned. The knee functioned satisfactorily for a further 15 years before requiring replacement.

Figure L.38 Osgood–Schlatter’s disease of the right knee. Note the prominence of the tibial tubercle secondary to the pull of the quadriceps muscle across the growth plate.
caused periosteal elevation. This bone destruction and periosteal reaction may not be radiologically evident for 10–12 days. In severe untreated cases, the pus strips the periosteum of the shaft and the nutrient artery becomes thrombosed, resulting in infarction of the diaphysis and eventual sequestrum formation. The periosteum continues to lay down new bone with the formation of an involucrum. The condition has now reached the chronic stage, and the clinical picture is completed by the presence of discharging sinuses. The disease is especially devastating if the child is malnourished, when there is a significant mortality due to either overwhelming sepsis or secondary abscess formation affecting in particular the brain, lungs and other bones. In modern practice, this fulminating picture is rarely seen, but if the diagnosis is delayed, the condition can still cause local destruction and long-term disability.

Distinguishing features between osteomyelitis and acute septic arthritis of the joint are: (i) in the latter, the synovial thickening, the effusion and the warmth are more marked; and (ii) the child will not even permit a jog of movement from the joint. Both infections are easily missed in neonates. Failure of the baby to spontaneously move a limb always requires an urgent diagnosis. The most common infecting organism of both osteomyelitis and septic arthritis is Staphylococcus aureus and due to MRSA the incidence has risen over the past decade. Streptococcal septic arthritis can be surprisingly persistent, even though the organism is sensitive to penicillin, as antibiotics do not penetrate well into the extensive fibrinous exudate that forms in the knee. Low-grade osteomyelitis or septic arthritis must be distinguished from the inflammatory arthropathies. Simple joint aspiration will confirm a septic arthritis while a MRI scan is good at localising the site of infection in osteomyelitis. In cases of doubt, a technetium scan or an Indium white cell-labelled scan can be helpful. Gonococcal arthritis is easy to overlook as it is a low-grade infection. The signs are not dissimilar from those of a mild haemarthrosis with an effusion, some synovial thickening and some warmth. If in any doubt about a knee effusion attributed by the patient to trauma, ask if there is a urethritis and look for a skin rash. Aspiration of the joint effusion will confirm the diagnosis. The diagnosis of tuberculous infection is frequently delayed, especially in Western medicine. Excessive quadriceps wasting is a feature of tubercular synovitis (Figs L.39 and L.40). The classic site for a Brodie’s abscess, which is a localized area of chronic osteomyelitis usually due to Staph. aureus, is in the proximal tibial metaphysis. It causes a dull, aching, persistent pain. The differential diagnosis is from an osteoid osteoma or a benign tumour.

**TUMOURS OF BONE**

**Malignant tumours**

Primary malignant tumours of bone are rare diseases. Metastases are far more common. The osteosarcoma, which is the most common of this rare group, affects children, adolescents and young adults. The majority present with pain, which is unremitting, usually worse at night and often initially attributed by the patient to an injury. In the early stages, the physical signs are very variable, and there may be little to find except some local tenderness. However, this is an aggressive, rapidly expanding tumour and a mass soon becomes evident, although the bony pathology may be camouflaged by the swollen overlying soft tissues.

**Figure L.39** Tuberculosis of the right knee. Marked muscle wasting of the thigh is a common finding, even in relatively early tuberculosis.

**Figure L.40** Same patient as Fig. L.39. Anteroposterior X-ray of both knees. There is marked generalized osteoporosis of the right knee, with ‘pencilling’ of the cortical margins and subchondral plates. Biopsy confirmed tuberculous synovitis.
There is often a small effusion within the knee, but movement is retained; this is the distinguishing feature from infection and myositis ossificans (see p. 360). X-ray features are variable, but essentially reveal destruction of the normal trabecular pattern, an indistinct margin and evidence of periosteal new bone formation – the classical ‘Codman’s triangle’ or ‘star-burst’ formation (Figs L.41 and L.42). A bone biopsy is required to establish the precise nature of the tumour.

Malignant transformation is a rare complication of Paget’s disease, but osteosarcoma secondary to Paget’s disease accounts for most osteosarcomas appearing in later life. This complication should be suspected in any long-standing Paget bone in which there is an alteration in the level of pain or swelling. The tumour is particularly aggressive, and many patients have pulmonary metastases by the time of diagnosis. Plain X-rays show areas of bone destruction and soft-tissue invasion, superimposed on the features of Paget’s.

Ewing’s sarcoma is the second most common primary malignant tumour of bone in children. It consists of islands of non-osteogenic and anaplastic small, round cells closely associated with blood vessels. The precise cell of origin remains unknown, but it is probably a neural cell. The patient is usually aged between 10 and 20 years, and presents with pain and swelling accompanied by warmth and tenderness, which, together with the systemic signs of pyrexia and a raised ESR, can mimic a low-grade osteomyelitis. X-rays show a permeating destructive lesion which, unlike an osteogenic sarcoma, is usually diaphyseal (Fig. L.43). The rapid expansion of the tumour produces periosteal elevation, which appears radiologically as a Codman’s triangle and ‘sun-ray’ spicules. Occasionally, this shows as the ‘onion-peel’ effect, due to repeated layers of periosteal bone being laid down around the diaphysis. CT and MRI imaging reveal a large extra-osseous component. Spread is to the lungs and other bones. The differential diagnosis is from primary lymphoma of bone and osteomyelitis.

Chondrosarcomas in particular arise from the flat bones, the pelvis and the scapula, often from a pre-existing exostosis (Fig. L.44). Consequently, any change in size in an exostosis in adult life needs urgent investigation. Chondrosarcomas are slow-growing lesions, and usually affect the 30-to 50-year age group.
Benign tumours

Giant-cell tumour of bone (osteoclastoma) derives its name from the presence of large, multinucleated giant cells. The tumour is usually considered benign but locally expansive within the bone. Although 2–10 per cent of these tumours metastasize to the lungs, the prognosis is still quite good, and far better than with other malignant bone tumours. Most patients are between 20 and 40 years of age. The most common site is the knee, followed by the sacrum and then the distal radius. The presenting feature is pain and swelling around the joint, which usually passes unnoticed for several months. The tumour has a unique classical appearance on X-ray as it involves the epiphyseal end of long bones and extends up to the joint margin (Fig. L.45). A delay in diagnosis increases the likelihood of a pathological fracture, which, if it occurs into the joint, makes subsequent treatment more difficult. Histologically, osteoclastoma is indistinguishable from the ‘brown tumour’ of hyperparathyroidism (see below p 378). With osteoclastoma the serum calcium is normal.

Osteochondroma (cartilage-capped exostosis) is the most common primary benign bone tumour. It has a bony base with a cartilage cap and occurs in the metaphyses of long bones, especially at the fast-growing joints (i.e. the knee, shoulder and wrist) (Fig. L.46). However, it can also be found in any bone of cartilaginous origin. It can be either solitary or multiple, and in the latter case may be hereditary (an autosomal dominant skeletal dysplasia). This lesion is generally considered to be a failure of remodelling at the growth plate and, when multiple lesions affect a single bone, this bone will be shorter as the growth potential of the bone has been partially squandered. The true incidence

Figure L.44 Chondrosarcoma of the left ilium (arrow) arising in a previous exostosis in a 35-year-old male who had noticed an increase in lesion size over the previous 6 months. There is a benign exostosis on the right ilium.

Figure L.45 Osteoclastoma (giant-cell tumour) in a 20-year-old female. The lesion extends up to the subchondral plate of the knee. Distally, the tumour margin is poorly defined, indicative of aggressive local invasion.

Figure L.46 (a) Cartilage-capped exostosis of the tibia. There is a well-defined prominence of the medial aspect of the proximal tibia. Anteroposterior X-ray showing the bony stalk (arrowed). The cartilage cap is radiolucent. (b) Lateral X-ray of the knee showing a large sessile exostosis of the distal femur.
is unknown, as many lesions are non-symptomatic and therefore undetected. Many are discovered purely as chance findings, often later in life. Most patients who present with either symptoms or a lump do so as teenagers or young adults. Osteochondromas are usually painless, but they can interfere mechanically with the function of tendons, which catch over the bony prominence. Since they are derived from the growth plate, growth of individual osteochondromata ceases when the patient stops growing. Any alteration of size later in life should raise suspicion of malignant transformation, usually to a chondrosarcoma (Fig. L.44.) Fortunately, this is rare, probably occurring in less than 1 per cent of solitary lesions, although more frequently, possibly up to 5 per cent, with multiple lesions.

Chondroblastoma is a benign tumour of cartilage and is distinguishable by being one of the few tumours to be confined to the epiphysis. It usually presents with a constant ache in the joint and localized tenderness in the adjacent affected bone. Patients are usually in their teens or early adult life (i.e. around the end of the growth period). X-rays classically show a rounded, well-demarcated radiolucent area in the epiphysis, often with an extension into the adjacent metaphysis (Fig. L.47).

Enchondromas also affect the metaphyseal–diaphyseal region and, like fibrous dysplasia, are centrally located. They are most frequently found in the metacarpals (Fig.L.48) or metatarsals, but they can occur in long bones, especially the humerus, femur and tibia. Radiologically in children, they show as well-defined cystic lesions, and it is not until the bone develops maturity that characteristic flecks of calcification show within the lucent area. Many are diagnosed incidentally, while some – particularly those in the hand – present with bony deformity. Do not dismiss pain as being due to a pathological fracture without good evidence, as malignant transformations to a chondrosarcoma can occur. This is rare in solitary lesions but more common in Ollier’s disease (multiple enchondromas) and Maffucci’s syndrome.

Chondromyxoid fibromas are rare benign tumours that also occupy the metaphyseal–diaphyseal junction. They are much more prevalent in the bones of the lower limb than the upper limb. They are eccentrically placed, and have a dense endosteal margin. They present with pain, and, in a superficial bone such as the tibia, there is overlying warmth and local swelling.

Tumour-like conditions
Many authorities consider that fibrous cortical defects and non-ossifying fibroma are essentially the same lesion. They are probably not true neoplasms, but rather...
developmental proliferations of fibrous tissue and histiocytes. Some 80 per cent of lesions occur in either the distal femur, the proximal tibia or the distal tibia. They are the most common bone lesion in children, being found in up to 30 per cent of children aged 4–8 years, and are usually discovered incidentally as they are asymptomatic (Fig. L.49). They may range from several millimetres to several centimetres in size. They heal spontaneously and do not require treatment unless there is a risk of a pathological fracture. Non-ossifying fibromas are merely larger and extend more deeply into the bone than the fibrous cortical defects (Fig. L.50). Consequently, there is a greater risk of pathological fracture.

Fibrous dysplasia arises in the marrow and is therefore situated centrally on a plain X-ray. It is composed of fibrous tissue and immature bone and occurs in two forms, solitary (80 per cent) or polyostotic (20 per cent). Common sites are the ribs, cranium, facial bones, femur (Fig. L.51), tibia and humerus. They are usually discovered incidentally on plain X-rays, on which they classically give an appearance of ‘ground glass’. However, appearances are very variable, and fibrous dysplasia is well-recognized as being able to mimic many other bone lesions. Most solitary lesions are asymptomatic, but they can present as a pathological fracture or bone deformity (Fig. L.52). The polyostotic form can be associated with a variety of endocrine disorders: hyperthyroidism, acromegaly, hyperparathyroidism, Cushing’s syndrome, diabetes and Albright’s syndrome – fibrous dysplasia, precocious puberty and pigmented skin lesions. Occasionally, the polyostotic form may undergo malignant transformation.

Osteoid osteoma can occur in any bone except the skull. It is a tiny, tumour-like condition that on excision appears as a dark brown or reddish nucleus surrounded by dense bone. Radiologically, lesions in the cortex show dense sclerosis and cortical thickening, which may mask the underlying small central nidus. This can be revealed by tomography (traditional technique), CT or MRI (Figs. L.53-55). A technetium bone scan is often helpful especially when the lesion is in the pelvis where the pain may be diffuse and the unfortunate patient labelled as a hysterical, psychotic or malingering. Now considered rather old fashioned, bone scans remain, in the author’s opinion, very useful in diagnosing rarities.

In a metaphysis an osteoid osteoma can be difficult to distinguish from a Brodie's abscess - a well-defined area of chronic osteomyelitis. In the spine, it is the classic cause of a painful scoliosis. The pain is immediately abolished by excising or ablating the lesion.

Osteoblastoma (giant osteoid osteoma) has similarities to an osteoid osteoma, although it is now officially classified as a benign tumour. The spine is the most
Figure L.51 Fibrous dysplasia. Anteroposterior X-ray of the distal femur, showing an extensive lesion affecting the medial half of the distal femoral metaphysis, resulting in inequality of growth.

Figure L.52 Same patient as Fig. L51. Clinically, there is a marked bowing of the left leg.

Figure L.53 Osteoid osteoma of distal tibia in a 17-year-old male. Although the patient gave a clear history of severe pain, well-controlled by simple analgesia, standard X-rays appeared normal.

Figure L.54 Same patient as Fig. L53. This bone scan showed an increase in uptake in the distal tibia, most marked posteriorly.

Figure L.55 Same patient as Fig. L53. Computed tomography scanning revealed the classical lesion of sclerosis with a small lucent central nidus in the posterior cortex of the tibia adjacent to the distal tibiofibular joint (arrowed).
common site. It is slow-growing and presents with mild persistent backache that is relieved by analgesics. Radiologically, the lesion is variable but is usually expansile, well-circumscribed and partially calcified. A simple bone cyst (unicameral) is a benign unilocular or partially locular fluid-filled cyst that occurs in the first two decades of life, most commonly in the proximal humerus and proximal femur. Many are found as chance findings on routine X-rays. These may never become symptomatic, and they appear to resolve spontaneously. Symptoms are the result of a pathological fracture through the weakened cortex (Fig. L.56). Plain X-rays are normally diagnostic as the cyst has a well-defined margin and never penetrates the cortex, which may be thinned and appear slightly expanded, particularly when the cyst migrates down from the metaphysis into the diaphysis. This expansion is never greater than the width of the overlying epiphyseal plate.

While aneurysmal bone cysts are classified as reactive lesions of bone (not neoplasms), they are nonetheless locally destructive. They can arise at any age and in any bone, but they are most commonly found in young adults affecting the metaphysis, especially the proximal humerus, the distal femur and the proximal tibia (Figs L.57 and L.58). They

Figure L.56 Unicameral bone cyst. There has been a pathological fracture through this cyst in an 8-year-old boy. The lesion has a well-defined margin and is located in the metaphysis, having 'migrated' from the epiphyseal plate.

Figure L.57 Aneurysmal bone cyst. Anteroposterior X-ray showing the expansion of the proximal fibula.

Figure L.58 Aneurysmal bone cyst. Operative specimen showing the typical vascular appearance of the lesion.
are blood-filled, expanding lesions and, although the overlying cortex is not destroyed, it may be rendered so thin as to be almost invisible on X-ray. Radiologically, they have a ‘soap bubble’ appearance due to bony ridges on the wall of the cyst. There are two types: primary, arising de novo; or secondary to bleeding into some other lesion, for example a giant-cell tumour. They usually present with discomfort, or occasionally as a bony swelling.

Osteitis fibrosa cystica (brown tumour) occurs in advanced hyperparathyroidism. Although uncommon, they nonetheless remain important diagnostically. Radiologically, they can be easily confused with other tumours and cysts of bone, while histologically they are indistinguishable from an osteoclastoma. (Pathologists will refuse to give an opinion on a biopsy from a lesion with a provisional diagnosis of osteoclastoma unless they have evidence that the calcium level is normal.) Multiple brown tumours are also easily confused with metastatic malignancy. Brown tumours are usually found in the long bones (Fig. L.59), the ribs or the skull (‘pepper-pot’ skull).

Other features include subperiosteal resorption in the phalanges, femoral neck, upper tibia and medial end of clavicle.

Apel’s diagram (Fig. L.60) is a useful aide memoire for this group of ‘cyst and cyst-like’ conditions of bone. Remember also that metastases are by far the most common malignant tumours found in bone, and that both metastatic and primary tumours can be confused with traumatic fractures (Fig. L.61).

**BURSITIS AND TENDINITIS**

Prepatellar bursitis (housemaid’s knee) is rarely seen in housemaids, but is an occupational hazard of those who do a lot of kneeling, for example carpet-layers. When uninfected, it presents as a smooth, circumscribed swelling anterior to the patella (Fig. L.62). Infrapatellar bursitis (clergyman’s knee) is more distally situated and lies over the patella tendon (Fig. L.63). When either superficial bursa becomes infected, the physical signs are obvious: tenderness, warmth and surrounding superficial oedema. The knee joint is not involved. Deep infrapatellar bursitis involves the bursa lying behind the patellar tendon, above its insertion onto the tibial tubercle. If this bursa is infected, the signs are less obvious than with the superficial bursae. The area of induration is more widespread, the tenderness is deeper and the erythema less marked. Flexion of the knee is uncomfortable and, as a consequence, tends to be restricted.

The semimembranosus bursa presents with a painless lump situated posteromedially; the bursa lies between the semimembranosus and the medial head of the gastrocnemius (Fig. L.64). It is most commonly seen in children and usually resolves spontaneously. The knee joint is normal. It needs to be distinguished from a popliteal cyst (Baker’s cyst). This is a herniation of the synovium, and is a sign of pathology within the knee (usually osteoarthritis),

![Figure L.59: Hyperparathyroidism (brown tumour).](image)

There has been a pathological fracture through this lesion in the proximal femoral diaphysis. Histologically, it is indistinguishable from an osteoclastoma.

![Figure L.60: Cyst and cyst-like lesions of bone.](image)
Figure L.61 (a) Pathological fracture of the distal humerus secondary to a renal carcinoma – hypernephroma. The initial X-rays were deceptive and the lesions considered to be a straightforward traumatic short spiral fracture. (b) An X-ray taken 6 weeks later shows the true destructive nature of the tumour, which is so vascular that, clinically, the lesion was pulsatile.

Figure L.62 Prepatellar bursitis. A prepatellar bursa lies directly over the patella.

Figure L.63 An infrapatellar bursa lies distal to the patella over the patellar tendon, before (upper) and after (lower) surgical excision.

Figure L.64 Semimembranosus bursa of the knee. This presents as a painless swelling in the popliteal fossa (arrowed), usually in children or adolescents. It is most prominent when the knee is straight.
although it can occur secondary to any pathology, including tuberculosis. Their importance lies in the fact that they may rupture and, when they do so, the signs are identical to those of a DVT, with a swollen tender calf. Patients need to be reassured as to the benign nature of the Baker’s cyst but warned about spontaneous rupture (Fig. L.65). Finally, do make certain that any cyst in the back of the knee is not pulsatile, i.e., a popliteal aneurysm!

**Meniscal cysts** arise from the meniscus–capsule junction, and are a reflection of abnormal mechanics within the meniscus, usually in association with a horizontal cleavage tear. Their pathology is identical to that of a ganglion of the wrist. The presentation between the two menisci is very different due to the lateral meniscal cyst being trapped under the dense iliotibial tract. **Lateral cysts** cause aching discomfort, present early when they are still small, and feel almost bony hard; indeed, they are frequently mistaken for an exostosis or an osteophyte. The lump is situated at or just below the level of the joint line, usually anterior to the collateral ligament, and is most evident with the knee slightly flexed. On full flexion, they often vanish in the sense that they are no longer palpable. **Medial meniscal cysts** are much less common. The signs are very different, as they are usually large, clearly cystic and relatively soft (Fig. L.66). Their peculiarity is that they frequently migrate a considerable distance away from the joint line and the meniscus to which they are attached by a ‘stalk’. They may be mistaken for a bursa that lies over the medial tibial condyle some 5 cm below the joint line at the insertion of sartorius. They can migrate into the fat pad anteriorly, or occasionally appear at the posteromedial corner of the knee, masquerading as a semimembranosus bursa in the wrong age group. A good-quality MRI usually reveals the meniscal attachment.

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**Figure L.65** (a) Transverse ultrasound scan of the popliteal fossa showing a Baker’s cyst surrounding the gastrocnemius muscle. (b) Arthrogram of the knee showing the extravasation of contrast into the calf from a ruptured Baker’s cyst.

**Figure L.66** Cyst of the medial meniscus of the right knee. (a) The cyst is lying over the medial tibial condyle, just below the joint line. These cysts can ‘migrate’ anteriorly into the fat pad, move further down the proximal tibia, or appear posteromedially in the popliteal fossa, as shown on the magnetic resonance imaging scan (b).
Tendinitides around the knee are common, as are muscular sprains. These may be due to acute injuries, chronic stress injuries, for example in middle-aged amateur marathon-runners, or secondary to an altered pattern of gait due to pathology elsewhere in the leg, for example malfunction of the quadriceps leading to overuse of the iliotibial band, with pain at the insertion onto the proximal anterolateral tibia.

PAIN IN THE CALF AND SHIN
Calf pain is a far more frequent symptom than pain affecting the anterior aspect of the lower leg. Furthermore, pain in the calf is commonly encountered with diseases that are either life-threatening or limb-threatening. Thus, calf pain requires a precise diagnosis and should never be dismissed as a muscular sprain.

Causes of calf or shin pain include:

**Life- or limb-threatening**
- DVT
- Arterial disease
  - Acute arterial ischaemia
  - Intermittent claudication
  - Compartment syndrome

**Less serious causes (see Box L.2)**
- Referred pain from the lumbar spine
  - Sciatica
  - Spinal stenosis
- Ruptured Baker’s cyst
- Muscle and tendon ruptures
  - Gastrocnemius/soleus ruptures
  - Ruptured Achilles tendon
- Varicose veins
- Nerve entrapment
  - Common peroneal nerve
  - Superficial peroneal nerve
- Peripheral neuropathies

**Life- or limb-threatening causes of calf pain**
The causes of the first group should never be missed. Both DVT and compartment syndromes can be very deceptive.

Deep vein thrombosis (DVT) requires permanent vigilance in most situations in which there is a known high incidence, especially postoperatively, bed rest for any illnesses, plaster splintage, air travel, etc. Postoperative prophylactic antithrombotic cover is no guarantee that this complication will not occur. Furthermore, those thrombi which are more likely to embolize often have less in the way of physical signs. For example, pulmonary embolism remains the main cause of death following routine hip replacement. Without prophylaxis, the incidence of DVT after major hip surgery is in the order of 60 per cent. This complication is not confined to surgical patients, but is also common in medical patients. After myocardial infarction or cerebrovascular accidents, the incidence has been estimated to be as high as 20 per cent. It is a well-described complication of the high-oestrogen contraceptive pill. It can even occur spontaneously in otherwise healthy individuals, for example on long-haul aircraft flights. The signs are usually unilateral. However, the apparently normal leg may be the site of a major embolus, since the signs become more apparent when the embolus becomes fixed in the vein. In terms of clinical features, look in particular for slight pitting oedema around the ankle; calf pain and tenderness; and a low-grade pyrexia. Homan’s test – pain on ankle dorsiflexion – is notoriously unreliable and should be discarded. In the ‘medical’ or outpatient setting, a low clinical probability score on the Wells criteria (Box L.4), plus a normal D-dimer assay, makes a diagnosis of venous thrombosis most unlikely.

In the early postoperative period, measurement of D-dimer – a fibrin degradation product – is elevated due to the surgical haematoma, and hence this assay is not diagnostic. Ultrasonic venous imaging is now the first line, but is less specific than radiological venography. Thus, if ultrasound is negative and clinical suspicion remains high, it should either be repeated 2 or 3 days later, or alternatively a venogram

**Box L.4 Wells clinical probability tool**

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<th>Wells explicit assessment</th>
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<tr>
<td>Active cancer</td>
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<td>Paralysis, paresis or recent plaster, or immobilization of lower limb</td>
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<td>Recently bed-ridden for more than 3 days, or major surgery in the last 4 weeks or more</td>
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<td>Localized tenderness</td>
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<td>Entire leg swollen</td>
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<td>Calf swelling &gt;3 cm compared with the asymptomatic leg</td>
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<td>Pitting oedema</td>
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<td>Collateral superficial veins</td>
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**Two-point deduction**
- Alternative diagnosis as likely or greater than deep vein thrombosis

Each positive response is 1 point, except if an alternative diagnosis is as likely as or greater than deep vein thrombosis, for which two points are deducted. 0 or fewer points low probability; 1–2 points moderate probability; 3 points = high probability.
should be performed. A massive proximal thrombosis affecting the iliofemoral veins can cause a very swollen white oedematous limb (phlegmasia alba dolens), but more commonly this presents as a blue leg (phlegmasia cerulea). An iliofemoral thrombosis can occasionally progress to true venous gangrene. The common long-term sequela of DVT is venous ulceration. It is now good practice that all patients admitted to hospital have a ‘UTE’ risk assessment. Intermittent claudication was discussed previously as part of the differential diagnosis of spinal stenosis (see ‘Leg pain of radicular or vascular origin’, p. 350).

Acute arterial ischaemia is classically due either to embolism – the most frequent cause is a thrombosis in the atrium secondary to atrial fibrillation – or to an acute thrombosis developing on a subintimal myocardial infarct. Other causes of acute ischaemia include an aneurysm (especially a popliteal aneurysm), an arteritis or trauma. Arterial injuries in association with fractures through the proximal third of the tibia are all too easy to miss, as the symptoms may be obscured by the major trauma to the limb. The effect of an embolus depends upon the extent and site of obstruction, the potential for collateral formation and the speed at which it becomes effective. Classically, emboli produce vasospasm with occlusion of the distal vessels leading to claudication, rest pain and gangrene. Clearly, if the occlusion is to a major vessel, the symptoms of pain are instant. Small peripheral emboli cause patches of cutaneous gangrene, with multiple small dark spots or blotches best seen in the foot.

Compartment syndrome is a deceptive and destructive condition due to muscle ischaemia as a result of a rise in pressure within the muscle compartment, eventually causing venous occlusion (Box L.5). A vicious circle is then set up in which capillary pressure rises towards arterial pressure, which results in diffusion of fluid into the extracellular compartment, causing a further rise in pressure. The more rigid the anatomical construction of the compartment, the more likely this syndrome is to occur. The forearm and the leg, which both contain two bones joined by a tough interosseous membrane and muscles contained within strong fascial compartments, account for the vast majority of cases. However, it can occur in other sites, for example the gluteal compartments, in which it can be particularly lethal due to a massive release of myoglobin, precipitating renal failure. An acute compartment syndrome is a surgical emergency and requires urgent decompression. Any delay over 4 hours leads to increasing disability secondary to muscle necrosis, resulting in fibrosis and contracture.

Soft-tissue crush injuries can be particularly deceptive, especially if the patient is seen shortly after the injury, before any swelling has become manifest. Never discharge a patient from the Accident and Emergency Department who has evidence of tyre marks across the calf (Fig. L.67)! Compartment syndrome also arises after prolonged crush injury, for example when a patient is trapped in a collapsed building or a motor car, and has sustained arterial compression and distal ischaemia.

Finally, in this group are included drug addicts who have been unconscious for long periods, resulting in local ischaemia. This is the most common cause of gluteal ischaemia. Open fractures of the tibia in which there has been wide soft-tissue destruction are less likely to suffer a compartment syndrome as the compartments have already been decompressed. The deceptive injury is the apparently straightforward

**Box L.5 Causes of compartment syndrome in the leg**

- **Acute traumatic**
  - Soft-tissue crush injuries
    - Acute severe local crush
    - Prolonged compression → ischaemia
    - Drug addicts secondary to coma
  - Fractures
  - Dislocation of knee
  - Haematoma

- **Postoperative**
  - Vascular
    - Post embolectomy
    - Post arterial reconstruction
  - Orthopaedic
    - Closed reduction of tibial fractures
    - Intramedullary nailing of tibia
    - Operations involving division of fibula (e.g. upper tibial osteotomy)
  - Bleeding disorders (e.g. haemophilia)

- **Chronic**
  - Muscle ischaemia in athletes

*Compartment syndrome is more common in closed than open injuries.

**Figure L.67** Extensive skin necrosis following a crush injury. The patient was run over by a car. In the Accident and Emergency Department, the only signs of the injury were tyre marks on the skin and a fractured fibula. One week later, the full extent of the injury is apparent.
closed fracture of the tibia and fibula in which the bones are overlapped. The chances of developing a compartment syndrome are increased by a closed reduction in which any overlap is fully corrected. The application of a plaster not only adds a further constriction to the limb, but also makes compartment syndromes more difficult to diagnose as the calf is not available to examine. However, treatment with intramedullary nailing also has a high risk – the process of reaming and putting down a nail produces a surge of pressure in the muscle compartments. Dislocation of the knee and fracture separation of the distal femoral epiphysis (an injury that occurs in the late teens before this epiphysis fuses) can both cause either lacerations or compression of the popliteal vessels, and consequently a high risk of calf compartment syndrome. Do not be deluded into believing that severe trauma is a prerequisite – the author has seen an anterolateral compartment syndrome following a kick at football causing a haematoma. In such circumstances, remember to check that the patient does not have a bleeding disorder. Compartment syndromes are well recognized following vascular surgery, especially post-embolectomy and after arterial reconstruction if there has been distal ischaemia for a significant period.

The diagnosis, as with DVT, is primarily dependent upon a high level of suspicion. The most important symptom is pain. It increases in severity to become unremitting. However, the pain will eventually subside and disappear when the muscle and the nerves are dead. The most important sign is pain on passive stretching of the involved muscle. This obviously involves a knowledge of which muscle is in which compartment (Table L.6). Due to the pain, the patient will be reluctant to move the limb. Clearly, the involved muscles will eventually be ‘paralysed’. In the early stages of the compression, paraesthesia can be present, but this is usually masked by the pain. The area of sensory loss is a very important sign, especially if the patient is in plaster and only the toes and the distal part of the foot can be examined. With regard to the pulse, beware! First, there may still be sufficient arterial pressure for the pulse to be transmitted through an involved compartment. Second, the pulse may have been transmitted via an unaffected normal compartment; for example, a compartment syndrome of the antero-lateral compartment will not obliterate the posterior tibial or the dorsalis pedis pulses. Finally, to complete the list of signs and symptoms commencing with the letter ‘P’, if the diagnosis is missed, the muscles will perish, the limb may perish, and so may the patient if this complication is further complicated by renal failure secondary to myoglobinemia.

In an established compartment syndrome, there is characteristic feel, described as ‘woody’. There are various methods of measuring intracompartmental pressure, but these can be unreliable. The old adage remains pertinent – ‘depend on the symptoms and the signs and, if in any doubt, decompress’ (Table L.7).

Other causes of calf pain
Referred pain from the lumbar spine, due either to radicular nerve involvement (sciatica) or to spinal stenosis, was discussed earlier (see ‘Leg pain of radicular or vascular origin’, p. 350). For a ruptured Baker’s cyst, see p. 380 and Fig. L.65. It is, of course, perfectly possible simultaneously to have a DVT and a Baker’s cyst, so proof must be obtained either that the cyst has ruptured and leaked or that the veins are patent. Muscle ruptures of either the gastrocnemius or the soleus occur at their attachment to the

<table>
<thead>
<tr>
<th>Table L.6 Compartments of the leg</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Compartment</strong></td>
</tr>
<tr>
<td>--------------------</td>
</tr>
</tbody>
</table>
| Anterior           | Tibialis anterior  
Extensor digitorum longus  
Extensor hallucis longus  
Peroneus tertius | Anterior tibial | Deep peroneal | Cleft between hallux and second toe |
| Anterolateral      | Peroneus longus  
Peroneus brevis | Peroneal | Superficial  
Peroneal | Dorsum of foot |
| Deep posterior     | Tibialis posterior  
Deep posterior  
Flexor hallucis longus | Posterior tibial | Tibial | Sole of foot |
| Superficial posterior | Gastrocnemius  
Soleus | Sural | Sural | Heel and lateral border of foot |
An Achilles tendon rupture normally occurs in older athletes, the 45- to 60-year age group, but it can occasionally be seen in early adult life. It is easily missed, since the 'gap' is obscured by swelling and the patient is still able to plantar flex the foot due to the action of other plantar flexors (tibialis posterior, etc.). They cannot, however, stand on tip-toe. The best sign is Simmond's test (Fig. L.68). Here, the patient lies prone with the feet over the end of the couch; first, observe the position of the foot – if the tendon is ruptured, the foot will be lying at a right angle. Then squeeze the calf – if the tendon is intact, the foot will move; if it is ruptured, the foot will not move. Peripheral neuropathies can be a cause of fleeting pains in the calf, so always beware of dual pathology. That is, if a nerve is irritable due to an underlying neurological disease, it will be much more prone to be symptomatic, or it will be paralysed by a superimposed mechanical disease such as disc or nerve root entrapment. Entrapments are uncommon in the leg, but they are well described for the common peroneal nerve at the level of the fibular neck and for the superficial peroneal nerve where it exits through the deep fascia at the junction of the middle and distal thirds of the leg, just anterior to the fibula. Gently tapping this point with the muscles under stress (resisted dorsiflexion and eversion of the ankle) causes paraesthesae. The common peroneal nerve is also at risk from a direct trauma, such as a kick at football and/or from inadequate protection during obstetric or urological surgery in the lithotomy position. The author has even seen a foot drop secondary to a very tightly applied crepe bandage.

### Shin pain

The common cause of pain over the anterior aspect of the proximal tibia is referred pain from the patellofemoral joint (see ‘Pain in the front of the knee’, p. 368). Shin splints are a complaint of runners. In the unfit who suddenly take up 'keep fit', there may be a stress fracture of the tibia. More commonly, there is a periosteal reaction along the lateral border of the tibial shaft. In both of these conditions, technetium bone scanning shows increased uptake (Fig. L.69). These bone changes must be distinguished from a chronic compartment syndrome secondary to muscle ischaemia. This particularly affects long-distance runners. There are other rare causes of pain such as the lightening pains of tabes dorsalis, while a periosteal reaction occurs in hypertrophic pulmonary osteoarthropathy. Paget's disease, in which there is increased bone turnover, causes tibial enlargement with anterior and lateral bowing (Fig. L.70). This can be painful early during the period of deformity when the bone is relatively soft. Pain in an established Paget’s is due to: (i) a stress fracture; (ii) development of osteoarthritis in the knee or ankle; or (iii) malignant transformation to an osteogenic sarcoma.

---

**Table L.7 Compartment syndrome: the cardinal symptoms and signs**

<table>
<thead>
<tr>
<th>Pain</th>
<th>The most important symptom</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pain on passive movement</td>
<td>The most important sign</td>
</tr>
<tr>
<td>Palpation</td>
<td>A solid 'woody feel' is a late sign. Urgent decompression is required but the muscles may already have suffered some permanent damage.</td>
</tr>
<tr>
<td>Paraesthesia/numbness</td>
<td>Useful in localizing which compartment</td>
</tr>
<tr>
<td>Paralysis</td>
<td>Can be a presenting feature of compartment syndrome in drug addicts following prolonged coma</td>
</tr>
<tr>
<td>Pulse</td>
<td>Very misleading (see text)</td>
</tr>
<tr>
<td>Pressure studies</td>
<td>Several methods. Useful in the unconscious patient, but requires experience both technically and with interpretation</td>
</tr>
</tbody>
</table>

N.B. If the clinical signs indicate compression, then decompress.
PAIN IN THE ANKLE AND HINDFOOT

A wide variety of injuries occur around the ankle, and in many instances these involve a combination of ligamentous and bony injury. They occur sequentially according to the severity of the injury. For example, in the common inversion injury, the anterior talofibular component of the lateral ligament is the initial structure at risk. In the simplest sprain, the tenderness is therefore located anteriorly on the lateral side of the talar neck, just in front of the ankle and over the anterior distal aspect of the fibula. Pain is aggravated by inversion of the foot at the subtalar joint. With increasing severity, the injury involves the rest of the lateral ligament complex, and marked swelling around the distal fibula is evident. As the force increases, a spiral fracture of the fibula results, and finally an additional shear fracture of the medial malleolus occurs as the talus disrupts the ankle mortice. Other patterns of injury result from adduction, abduction and eversion injuries. The important initial assessment is to establish whether or not the talus is stable in the ankle mortice. Beware of the ankle that has marked tenderness over both malleoli, even if the X-ray shows only an undisplaced fracture of one of the malleoli, as a fracture on one side and a ligamentous injury on the other side equals ‘potential instability’. If in doubt, take stress X-rays of both ankles for comparison.

**Box L.6** Brief summary of causes of ankle pain

<table>
<thead>
<tr>
<th>Traumatic</th>
<th>Infective</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fractures (often referred to as Pott’s fractures)</td>
<td>Septic arthritis</td>
</tr>
<tr>
<td>Ligament sprains/ruptures</td>
<td>Osteomyelitis</td>
</tr>
<tr>
<td>Tendinitis: acute, chronic, rupture</td>
<td></td>
</tr>
<tr>
<td>Osteochondritis dissecans</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Inflammatory</th>
<th>Referred pain</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rheumatoid arthritis</td>
<td>Lumbar disc disease with ‘sciatica’</td>
</tr>
<tr>
<td>Degenerative</td>
<td></td>
</tr>
<tr>
<td>Osteoarthritis</td>
<td></td>
</tr>
</tbody>
</table>

**Figure L.69** Shin splints in a 20-year-old athlete. The technetium scan shows an increased uptake in both tibial diaphyses, more marked on the right than the left.

**Figure L.70** Paget’s disease of the right tibia. Note the typical varus deformity.
Tendinitis, whether acute or chronic, is common around the ankle. The tendons involved are:

- Posteriorly: Achilles tendon
- Laterally: peroneus longus/brevis
- Medially: tibialis posterior, flexor hallucis longus and flexor digitorum longus
- Anteriorly: tibialis anterior, extensor hallucis longus, extensor digitorum longus and peroneus tertius

A painful limp is the usual presenting feature, while the signs are the classical triad of local tenderness, pain on active resisted movement and pain on passive stretching. Note whether there is marked swelling along the involved tendon sheath as this is often indicative of incipient tendon rupture. A common example is a rupture of the tibialis posterior tendon, which leads to a classical deformity with marked valgus at the heel, a fallen longitudinal arch and a flat, externally rotated foot (Fig. L.71). A tender, swollen tibialis tendon sheath is an important sign as this pre-rupture stage often responds to treatment with simple orthotics - a heel cup with a medial longitudinal arch support.

Osteochondritis dissecans affects the dome of the talus, usually at the superomedial corner of the articular surface. This presents with aching discomfort aggravated by use, or occasionally with locking if the fragment has separated. The fragment is often small and can be missed on routine X-rays if their quality is poor.

Septic arthritis can be difficult to diagnosis if it is low grade or if it occurs as a complication of rheumatoid arthritis. Tuberculous infection can occur in the ankle joint, but it may also affect a tendon sheath. Synovial thickening and warmth from the ankle is best felt behind the malleoli, the normal concavity on either side of the Achilles tendon being obliterated (Fig. L.72). The distal tibial metaphysis and metatarsal bones of the feet are common sites for osteomyelitis. Since the bones are superficial, signs of inflammation are often present in the overlying skin. It is therefore easy to confuse osteomyelitis with cellulitis, indeed, so much so that the old rule states, ‘a cellulitis over a superficial bone is an osteomyelitis until proven otherwise’. Both the ankle and the subtalar joints are frequently involved in rheumatoid disease. This produces a characteristic valgus hindfoot deformity, which is occasionally complicated by a tarsal tunnel syndrome with compression of the plantar nerves. Any severe valgus deformity can cause localized pain at the distal end of the fibula due to impingement of this bone on the calcaneus, and consequent entrapment of the peroneal tendons. An acute exacerbation of pain in the hindfoot, in the absence of a flare of rheumatoid in other involved joints, should raise suspicion of a superimposed septic arthritis.

Significant symptoms due to osteoarthritis of the ankle are uncommon – a surprising fact considering how frequently this joint is injured. However, since the joint works in its mid-range, it rarely becomes

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**Figure L.71** Flat foot secondary to rupture of the left tibialis posterior tendon. The heel is in valgus, the medial longitudinal arch has collapsed, and the foot is pronated. Viewed from behind, the toes of the left foot are visible – the ‘too many toes’ sign.

**Figure L.72** Tuberculous synovitis of the right ankle in a 16-year-old boy. Note the loss of concavity on both sides of the right Achilles tendon (arrowed).
symptomatic until there has been sufficient loss of mobility to cause a fixed equinus deformity. The precipitating cause is usually obvious – malunited fracture, severe ligamentous instability, osteochondritis dissecans of the talus, previous infection, inflammatory arthropathies or bleeding disorders. Similar comments apply to osteoarthritis of the subtalar joint. The most common cause is incongruity secondary to fractures of the calcaneus. The natural history following this injury is for the pain gradually to settle over a period of 18 months to 2 years, but the joint remains stiff and the heel remains widened; this can cause problems with shoe-fitting. However, it is surprising how well patients adjust to this disability given sufficient time.

Several entities present specifically with pain below or behind the heel. These are age-related. Sever’s disease is a traction apophysitis of the Achilles tendon that most commonly affects boys aged 10–12 years, who present with a transient discomfort and mild tenderness localized to the posterior aspect of the calcaneus (Fig. L.73). ‘Heel-nobs’ is a complaint of teenage girls, the prominence on the back of the calcaneus being aggravated by fashionable shoes. Tight-fitting shoes or boots in young adults can result in an Achilles bursitis localized just above the insertion on the calcaneus. Achilles peritendonitis is a common problem in athletes of all ages, but it can also be precipitated by ill-fitting shoes. Plantar fasciitis occurs acutely in association with Reiter’s disease or ankylosing spondylitis in young men. Occasionally, it may be the presenting symptom of these diseases. The common type of plantar fascitis, ‘policeman’s heel’, occurs in 40- to 60-year-olds. There are no signs of overt inflammation, and the pain and tenderness are localized under the heel. Symptoms are usually worst first thing in the morning, and the first few steps taken on getting out of bed are particularly uncomfortable. If symptoms persist, check the uric acid level for gout. A bony spur is sometimes present on the lateral X-ray of the calcaneus, but this is also present in many patients who never have any symptoms of heel pain. A better indication for an X-ray is to exclude diseases of the calcaneus, for example tuberculous osteomyelitis or Paget’s disease, which can masquerade as a plantar fasciitis.

PAIN IN THE MID-FOOT (MID-TARSUS)
See Table L.8.

Kohler’s disease (in children) (Fig. L.74) and Brailsford’s disease (middle-age in women) both affect the navicular, and are both best regarded as an avascular necrosis. A bony prominence over the dorsal surface of the medial cuneiform and adjacent first metatarsal (a bone knob) is a common condition in the high-arch foot and can cause pressure against the shoe. Tibialis posterior and tibialis anterior tendinitis with tenderness and discomfort localized to the insertions on the medial side of the mid-foot is another common condition often associated with ‘falling arches’; there is a secondary planovalgus mid-foot deformity. Tenderness at the base of the fifth metatarsal is usually precipitated by an inversion sprain, as both the peroneus brevis and

Figure L.73 Sever’s disease: osteochondritis of the epiphysis of the calcaneus. Although both X-rays show similar fragmentation, only the right foot was painful.

<p>| Table L.8 Summary of causes of pain in the mid-foot |</p>
<table>
<thead>
<tr>
<th>Condition</th>
<th>Site</th>
<th>Diagnostic entity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Osteochondritis/</td>
<td>Navicular</td>
<td>Children: Kohler’s disease</td>
</tr>
<tr>
<td>Osteonecrosis</td>
<td>NAV</td>
<td>Adults: Brailsford’s disease</td>
</tr>
<tr>
<td>Bone overgrowth</td>
<td>Medial cuneiform/first metatarsal</td>
<td>Bone knob</td>
</tr>
<tr>
<td>Tendinitis</td>
<td>Navicular tuberosity</td>
<td>Tibialis posterior</td>
</tr>
<tr>
<td></td>
<td>Medial cuneiform/first metatarsal</td>
<td>Tibialis anterior</td>
</tr>
<tr>
<td></td>
<td>Base of fifth metatarsal</td>
<td>Peroneus brevis</td>
</tr>
<tr>
<td>Osteoarthritis</td>
<td>Talonavicular joint</td>
<td>Diabetes</td>
</tr>
<tr>
<td></td>
<td>Calcaneocuboid joint</td>
<td>Other peripheral neuropathies</td>
</tr>
<tr>
<td></td>
<td>Cuneiform–metatarsal (1st, 2nd, 3rd)</td>
<td>Tabes dorsalis</td>
</tr>
<tr>
<td></td>
<td>joints</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cuboid–metatarsal (4th, 5th) joints</td>
<td></td>
</tr>
<tr>
<td>Rheumatoid arthritis</td>
<td>Pan-tarsal involvement</td>
<td></td>
</tr>
<tr>
<td>Charcot joint</td>
<td>Pan-tarsal involvement</td>
<td></td>
</tr>
<tr>
<td></td>
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</tr>
</tbody>
</table>
tertius tendons attach here. Quite frequently, there is an undisplaced avulsion fracture at the base of this bone, since normal tendons are stronger than bone. Mid-foot osteoarthritis is common in middle years and later life, and the symptoms are usually mild. Charcot neuropathic joints are often initiated by a stress fracture occurring into the articular surface, and at this stage they may be painful. Rapid destruction of the mid-foot soon occurs. The foot shortens and the joints subluxate to produce the classical deformity of the ‘square tarsus’ (Fig. L.75).

PAINFUL FOREFOOT (METATARSALGIA)
The causes of a painful forefoot include:

Conditions affecting the hallux:
- Hallux rigidus
- Hallux valgus/bunion
- Rheumatoid arthritis/gout

Conditions affecting the metatarsals (two, three, four and five):
- Freiberg’s disease (an osteonecrosis of the second metatarsal head)
- Kohler’s second disease (an osteonecrosis of the third metatarsal head)

Figure L.74 Osteonecrosis of the navicular (arrow): Kohler’s disease.

Figure L.75 Anteroposterior (a) and lateral (b) X-rays of a Charcot foot in a diabetic patient. There is dislocation and disintegration across the mid tarsus (arrow).

- Antebunion (a bursa over a prominent fifth metatarsal head)
- Stress fracture of the metatarsal shaft
- Morton’s neuroma

Conditions affecting the metatarsophalangeal joints:
- Rheumatoid arthritis
- Traumatic dislocation/subluxation

Other conditions:
- Plantar warts
- Peripheral neuropathy
- Erythromelalgia
- Ischaemia
- Ingrowing toenails
- Subungual exostosis
- Onchogryphosis

Hallux rigidus is a stiff great toe secondary to osteoarthritis of the first metatarsophalangeal joint. Pain is caused by the loss of movement preventing the passive hyperextension of the joint, which is required for ‘toe-off’ at the end of the normal gait cycle.
The toe is usually in good alignment, but there is bony swelling around the joint secondary to osteophyte formation. Plantar flexion and dorsiflexion are both markedly restricted. The pain is best reproduced by moving the joint while compressing the toe against the metatarsal. Many cases of hallux valgus are painless, the metatarsophalangeal joint retaining a good range of movement. Symptoms are usually due to wearing shoes that can no longer accommodate the widened splayed forefoot – the first metatarsal moves medially, while the toe itself moves valgus (i.e. laterally), and as it does so the lesser toes are also displaced in a valgus direction or else become overriding (Fig. L.76).

A bunion is due to an adventitial bursa forming over the medial side of the prominent metatarsal head. This is prone to secondary infection. The severe pain of acute gouty arthritis accompanied by the surrounding erythema and oedema is very characteristic (Fig. L.77). Rheumatoid disease commonly affects the metatarsophalangeal joints, usually in a symmetrical pattern in both feet.

Pain and local tenderness under the metatarsal heads is a common ‘mechanical’ problem, especially in a cavus foot with curly toes, a combination that drives the metatarsal heads into the sole of the shoe. During the gait cycle in the normal foot, the pressure zone moves from the heel along the lateral border of the foot, and then swings back medially across the metatarsal heads to the hallux. Any bony prominence will create a high-pressure point resulting in thickening of the overlying skin, thus increasing the effect of the bony prominence and thereby establishing a vicious circle, resulting in a corn or a callosity. The pressure pattern can be studied using a pedobarograph, but sufficient diagnostic information can simply be found in the clinic by studying the pattern of callosities, the wear on the sole of the shoe, and the presence of humps and hollows in the insole.

Freiberg’s and Kohler’s second disease (Fig. L.78) are best regarded as an osteonecrosis that affects the second and third metatarsal heads, respectively. Stress fractures can affect any metatarsal, but most commonly occur in the necks of the second and third, usually following a prolonged period of unaccustomed activity (Fig. L.79). Morton’s neuroma is a swelling of digital nerve secondary to entrapment between the metatarsal heads. Pain is aggravated by walking and relieved by rest, particularly by removing the shoe. It is sometimes accompanied by paraesthesia of the appropriate toes. On examination, there are frequently no abnormal findings. The most useful sign is to apply cross-pressure to the metatarsal heads by squeezing the forefoot transversely while at the same time, with the other hand, eliciting the tenderness localized to the affected cleft; the third/fourth cleft is the most commonly involved. Demonstrable loss of sensation of the adjacent sides of the toes of the affected cleft is confirmatory, but is unfortunately an infrequent finding.

A bunion, termed an antebunion, can also form over a prominent fifth metatarsal head, often in association with a hallux valgus deformity consequent to widening of the forefoot and its entrapment in a shoe.
Plantar warts (verrucas) are particularly troublesome if they occur directly beneath a metatarsal head. Superficially, they can be difficult to distinguish from a corn, but paring down the skin with a scalpel blade will reveal the characteristic circular pattern of the verruca. Bone and joint infections in normal feet are uncommon unless there has been a penetrating injury. However, it is all too easy to overlook bone infection in an ulcerated neuropathic foot, especially in diabetes. Immunocompromised patients and those with HIV are also at risk. Peripheral neuropathy, particularly in the early stages, can cause burning pain in the foot. The common causes are diabetes, alcoholism, nutritional deficiencies (especially vitamin B) and HIV infection. Burning pain also occurs in the unusual condition of erythromelalgia. This is an abnormality of the superficial blood vessels found in polycythaemia, syringomyelia or the early stages of chronic arsenical poisoning. The pain is followed by cutaneous flushing and cyanosis, with the pain becoming pulsatile in character. Oedema, tenderness and hyperhydrosis may all be present. Occasionally, Raynaud’s phenomenon can produce similar episodic pain. Rest pain that worsens at night is a classical symptom of critical limb ischaemia secondary to peripheral vascular disease. The pain is dull, severe and often requires opiate analgesics. Classically, the patient is woken with a painful cold foot that is relieved by hanging the foot over the side of the bed or walking about. In critical ischaemia, pressure at the ankle is less than 50 mmHg, or 20 mmHg at the toe in patients who are not diabetic. Signs of ischaemia vary from dry necrosis of the skin on the toes to gangrene. Nailfold infection can be a presenting sign of arterial insufficiency. Do not confuse this with the infected ingrowing toenail of the hallux that is most frequently seen in teenage boys (Fig. L.80).

A subungual exostosis (Fig. L.81) arises from the terminal phalanx of the hallux, and the progressive enlargement causes deformity of the nail and pain secondary to pressure from the nail and the shoe. An onychogryphosis (Fig. L.82) is a disordered growth of the nail, usually secondary to trauma to the nail bed. The nail fails to grow forward, becomes thickened and deformed, catches against the shoe, and is difficult to cut.

DEFORMITIES OF THE FOOT

See Box L.7.

The term ‘talipes’ denotes a foot deformity affecting the hindfoot; varus describes the inverted heel; valgus the everted heel; equinus the plantar-flexed foot; and calcaneus the dorsiflexed foot. Mid-foot deformities are pes cavus, the high arch foot, and
pes planus, the flat foot. The metatarsals are referred to as adducted or varus, in which the metatarsal points medially, while metatarsus valgus means that the metatarsal points laterally and is an uncommon deformity usually due to a surgical overcorrection of a metatarsal osteotomy for hallux valgus. If the foot is supinated, it is inverted and rotated so that the first metatarsal tends to rise off the ground, the longitudinal arch is accentuated and weightbearing occurs along the lateral border of the foot. A pronated foot is the reverse, the foot appearing flat with depression of the longitudinal arch, and the first metatarsal head being forced into the ground.

With any foot deformity in childhood, it is essential to examine the peripheral neurology and also the spine. Talipes equinovarus (club foot) (Fig. L.83) describes a combined deformity of plantar flexion and inversion of the hindfoot, usually accompanied by a metatarsus adductus deformity arising at the tarsometatarsal joint. It is classified into three groups: (i) postural (due to intrauterine position, for example in oligohydramnios); (ii) idiopathic, in which the child is otherwise normal; and (iii) syndromic (acquired), in which there are other generalized abnormalities (e.g. arthrogryposis or diastrophic dwarfism) or neurological imbalance (e.g. myelomeningocele or spinal dysraphism).

Club foot occurs approximately 1 per 1000 live births. Males are twice as commonly affected as females, but although there is a familial incidence with an increase in first-degree relatives, the precise genetics remain unknown. One-third of cases are bilateral. Formerly, these were classified into mobile, semi-correctable and uncorrectable. This classification has largely been discarded because it is of little help in guiding treatment. The introduction of the Ponceti method of serial casting combined with manipulation, has greatly reduced the requirement for operation. Of all idiopathic (congenital) club feet treated by the Ponceti method about 70% will require a percutaneous Achilles tendon tenotomy and of that group, a further 10% will require revision.

Box L.7 Summary of paralytic causes of foot deformities*

<table>
<thead>
<tr>
<th>Upper motor neurone lesions</th>
<th>Lower motor neurone lesions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cerebral</td>
<td>Amyotrophic lateral sclerosis</td>
</tr>
<tr>
<td>Cerebral palsy</td>
<td>Cauda equina lesions (e.g. tumours, diastematomyelia)</td>
</tr>
<tr>
<td>Meningitis</td>
<td>Poliomyelitis</td>
</tr>
<tr>
<td>Cerebrovascular accident</td>
<td></td>
</tr>
<tr>
<td>Spinal</td>
<td></td>
</tr>
<tr>
<td>Trauma</td>
<td></td>
</tr>
<tr>
<td>Haemorrhage</td>
<td></td>
</tr>
<tr>
<td>Thrombosis</td>
<td></td>
</tr>
<tr>
<td>Spinal dysraphism</td>
<td></td>
</tr>
<tr>
<td>Diastematomyelia</td>
<td></td>
</tr>
<tr>
<td>Lower motor neurone lesions</td>
<td></td>
</tr>
<tr>
<td>Peroneal muscular atrophy</td>
<td></td>
</tr>
<tr>
<td>(Charcot–Marie–Tooth, which is now classified as an inherited mixed sensorimotor neuropathy)</td>
<td></td>
</tr>
<tr>
<td>Progressive muscular atrophy</td>
<td></td>
</tr>
<tr>
<td>(motor neurone disease)</td>
<td></td>
</tr>
</tbody>
</table>

*Certainment. In all foot deformities, examine the spine and the neurology.

The first metatarsal tends to rise off the ground, the longitudinal arch is accentuated and weightbearing occurs along the lateral border of the foot. A pronated foot is the reverse, the foot appearing flat with depression of the longitudinal arch, and the first metatarsal head being forced into the ground. With any foot deformity in childhood, it is essential to examine the peripheral neurology and also the spine.

Talipes equinovarus (club foot) (Fig. L.83) describes a combined deformity of plantar flexion and inversion of the hindfoot, usually accompanied by a metatarsus adductus deformity arising at the tarsometatarsal joint.

* N.B. In all foot deformities, examine the spine and the neurology.

Figure L.80 Infected ingrowing toenails in a teenage boy.

Figure L.81 Subungual exostosis. Left: Preoperatively. Right: At surgery, the exostosis is a well-defined spherical nodule attached to the distal phalanx.

Figure L.82 Onychogryphosis of the toenails of both big toes.
percutable tenotomy. A further small percentage of these babies will need later secondary intervention either with postero-medial release or tibialis anterior transfer.

_Talipes calcaneovalgus_ is a common mild deformity secondary to intrauterine moulding. It corrects easily with manual stretching, and the foot becomes normal.

Adduction deformities of the forefoot occur in association with a congenital club foot, but they also arise independently; this is _congenital metatarsus varus_ (Fig. L.84). This is a common deformity in the baby and young child. The vast majority resolve spontaneously with growth, and surgical correction is rarely required.

_Pes planus_ (sometimes referred to as pes valgus as the heel is a valgus position) describes the flat foot. This is a common, indeed almost universal, finding in young children, the normal arch only gradually appearing over the first few years. Ensure, however, that the foot is mobile.

_Congenital vertical talus_ is a rare disorder in which there is a talonavicular dislocation with the navicular lying on top of the neck of the talus (Fig. L.85). The foot is stiff and consequently acts in a ‘rocker-bottom’ manner. Occasionally, a flat foot persists into adult life, when it may be associated with other causes of ligamentous laxity, for example Ehlers-Danlos syndrome or Marfan’s syndrome. _Spasmodic flat foot_ is a well-recognized entity, although the foot is not necessarily flat, nor is the problem necessarily spasmodic. This condition is due to a bony union of the tarsal bones, a talocalcaneal bar being the most common type. The foot is held rigidly everted secondary to muscle spasm. The peroneal and extensor tendons stand out prominently (Figs L.86 and L.87). Subtalar movements are grossly restricted and often painful. Mid-tarsal movements are diminished, but ankle movements are usually normal. Special oblique X-ray views are required to demonstrate a bar or an abnormal joint. Technetium bone scanning can be helpful in pinpointing the site of the anatomical problem, which is usually best shown by a CT scan. Inflammatory conditions affecting the subtalar–mid-tarsal joints can also cause this deformity. A pes planus is common in _paralytic disorders_. In the elderly, it can be associated with a general muscular weakness.
aggravated by obesity, while a more specific cause is rupture of the tibialis posterior tendon (see ‘Pain in the ankle and hindfoot’, p. 385).

The majority are ‘idiopathic’, but ensure that there is no underlying neurological or muscular deficit causing a muscle imbalance that particularly affects the intrinsic muscles. Charcot–Marie–Tooth (peroneal muscular atrophy) (Fig. L.88), Friedreich’s ataxia and spastic hemiplegia are some examples. Pes cavus was a common deformity in polio (Fig. L.89). In general, pes cavus is a much more troublesome entity than pes planus, since weightbearing tends to be concentrated

(a)

(b)

Figure L.86 Spasmotic flat foot. Upper: The peroneal tendons stand out sharply in spasm. Lower: The examiner is unable to invert the hindfoot, which is held rigid.

Figure L.87 Same patient as in Fig. L.88. An X-ray confirms the presence of a bony bar between the navicular bone and the calcaneus.

A pes cavus deformity is usually associated with clawing of the toes in addition to the high arch.

Figure L.88 Hereditary motor sensory peripheral neuropathy (Charcot–Marie–Tooth syndrome). The foot is cavus, the toes are clawed (a), and there is marked wasting of the distal calf (b).

Figure L.89 A severe pes cavus deformity secondary to a neurological imbalance, in this case due to poliomyelitis.
along the lateral border of the foot and on the metatarsal heads. In contrast, in the flat foot, weight is spread over a wide area. Interestingly, even when there is an underlying pathological reason for a pes cavus, the deformity is rarely seen in children under the age of 6 years.

DEFORMITIES OF THE TOES

The fundamental problem in the deformities of claw-toe, hammer toe and cock-up toe is now considered to be the dorsiflexive contracture at the metatarsal-phalangeal joint. In the claw toe deformity, there is in addition to the hyperextension of the metatarsal-phalangeal joint flexion contractures of the interphalangeal joints. It is often associated with a pes cavus deformity. Initially, the toe deformities are mobile and can be passively corrected, but they gradually become fixed, and eventually the metatarsophalangeal joints may subluxate or even dislocate. The condition is usually bilateral. While most cases are idiopathic, it can result from a peripheral neuropathy; in particular, peroneal muscular atrophy should be excluded. Claw toes are also commonly found in rheumatoid arthritis secondary to subluxation and dislocation of the metatarsophalangeal joints. Symptoms are secondary to pressure on the prominent metatarsal heads and also over the dorsum of the toes. Callosities are usually present at both sites. In severe cases, walking may be difficult.

In a hammer toe deformity, the proximal interphalangeal joint is fixed in acute flexion while the metatarsophalangeal and distal interphalangeal joints are fixed in extension. The second toe is the most commonly affected. Symptoms are due to painful callusities forming under the metatarsal head and over the dorsum of the toe. In a cock-up toe deformity, the metatarsophalangeal joint is dislocated, with the result that the little toe contracts proximally and sits on the dorsum of the metatarsal head. An overlapping little toe is a common anomaly in which the little toe sits on the dorsum of the fourth toe. Both little toe deformities can cause problems with shoe-fitting. In a mallet toe deformity, there is fixed flexion of the distal interphalangeal joint so that the tip of the toe or the toenail is impacted down into the sole of the shoe, resulting in a painful callosity.

LYMPHADENOPATHY

Mark Kinirons

Although there are a multitude of causes of lymphadenopathy (Box L.8), it is usually possible to narrow the differential diagnosis considerably by a careful review of the patient's history and clinical examination. The differential diagnosis is very dependent on the patient's age and geographical location. Small soft lymphadenopathy is commonly found in healthy children and young adults, particularly in the axillary and inguinal regions. A young adult with significant cervical lymphadenopathy in the UK is likely to have Epstein–Barr virus (EBV) infection, whereas in Africa the most common cause is often tuberculosis. Local lymphadenopathy may be secondary to local infection or malignancy, whereas generalized enlargement is indicative of a systemic disorder. Painful, tender nodes in association with a fever and rash indicate a systemic infection, usually of viral aetiology. Firm, painless nodes found in a patient with weight loss or night sweats are likely to be due to malignant disease. Occasionally, it is difficult to identify the cause of enlarged lymph nodes, and it is then essential to systematically consider all possibilities listed in Box L.8.

INFECTIONS

Local infections usually result only in regional lymph node enlargement, which may be painful. A viral sore throat may cause swelling of the tonsils and tonsillar lymph nodes. Unilateral cervical, axillary or inguinal

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### Box L.8 Causes of lymphadenopathy

<table>
<thead>
<tr>
<th>Infectious diseases</th>
<th>Malignant diseases</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Viral</strong>: Epstein–Barr virus, cytomegalovirus, HIV, Caxsackie virus, hepatitis viruses, rubella, measles, herpes simplex, varicella-zoster</td>
<td><strong>Secondary deposits from a local malignancy</strong></td>
</tr>
<tr>
<td><strong>Bacterial</strong>: Local infections (Staphylococcus, Streptococcus), brucellosis, tuberculosis, atypical mycobacteria, syphilis, cat scratch disease, plague, leprosy, bejel, pinta, yaws, bartonellosis, tularaemia</td>
<td><strong>Hodgkin's disease</strong></td>
</tr>
<tr>
<td><strong>Protozoan</strong>: Toxoplasmosis, trypanosomiasis, leishmaniasis, filariasis</td>
<td><strong>Non-Hodgkin's lymphomas</strong>: B-cell, T-cell, mycosis fungoides, angioimmunoblastic lymphadenopathy</td>
</tr>
<tr>
<td><strong>Fungal</strong>: Blastomycosis, histoplasmosis, paracoccidiomycosis</td>
<td><strong>Acute or chronic lymphatic leukaemia</strong></td>
</tr>
<tr>
<td><strong>Chlamydial</strong>: Lymphogranuloma venereum, trachoma</td>
<td><strong>Myeloproliferative disorders, macroglutulinaemia, amyloidosis</strong></td>
</tr>
<tr>
<td><strong>Rickettsial</strong>: Scrub typhus</td>
<td><strong>Immunological diseases</strong>: Systemic lupus erythematosus, Rheumatoid arthritis, juvenile rheumatoid arthritis</td>
</tr>
<tr>
<td><strong>Protozoan</strong>: Toxoplasmosis, trypanosomiasis, leishmaniasis, filariasis</td>
<td><strong>Mixed connective tissue disease</strong></td>
</tr>
<tr>
<td><strong>Drug reactions</strong>: Phenyoitin, hydralazine, gold, allopurinol</td>
<td><strong>Serum sickness</strong></td>
</tr>
<tr>
<td><strong>Miscellaneous</strong>: Sarcoïd</td>
<td><strong>Amyloidosis</strong></td>
</tr>
<tr>
<td><strong>Lipid storage diseases</strong></td>
<td></td>
</tr>
</tbody>
</table>
lymphadenopathy may be due to bacterial or fungal infection in the field of lymphatic drainage.

*EBV infection* (glandular fever) is a common cause in young adults with generalized lymphadenopathy particularly involving the cervical region. It is clinically very similar to acute cytomegalovirus (CMV) and HIV infection. This latter virus may later cause a persistent, painless, generalized lymphadenopathy for a period of many months. *Acute brucellosis* or *toxoplasmosis* may also produce a similar syndrome, although usually without a sore throat.

*Tuberculosis* usually causes enlargement of a single set of nodes, although generalized enlargement is occasionally observed. *Primary syphilis* may initially result in swelling of the inguinal lymph nodes draining the chancre, but generalized lymphadenopathy may be observed in *secondary syphilis* when the primary lesion has healed. *Cat scratch fever*, which is often seen in children, presents with an inflamed papule at the site of injury, and this may be followed by local lymphadenopathy for several months.

*Tropical infections* such as filariasis, African trypanosomiasis and tularaemia may be associated with lymphangitis in association with the primary infection, and local lymphadenopathy that may later become generalized.

**MALIGNANT DISEASE**

Malignant disease may spread from a primary site along the draining lymphatics to invade the regional lymph nodes; it is unusual for generalized lymphadenopathy to be secondary to disseminated carcinoma, although it is occasionally observed (e.g. melanoma). *Virchow's node* is found in the anterior left supraclavicular region, and is also known as Trousier's ganglion. Causes include carcinoma of the breast or bronchus, gastrointestinal malignancy and lymphomas.

Symmetrical generalized painless lymphadenopathy is a common presenting feature of malignancies of the lymphopoietic system – for example, Hodgkin's disease in a young person, or chronic lymphatic leukaemia or non-Hodgkin's lymphoma in an older adult. *Macroglobulinaemia*, or occasionally *myeloma*, is characterized by painless generalized lymphadenopathy. Such syndromes may be associated with systemic or 'B' symptoms – i.e. over 10 per cent weight loss, fever of over 38 °C or night sweats. These indicate generalized disease, and in an individual presenting with only a single group of enlarged nodes, they are suggestive of more extensive disease. Other malignant disorders of the lymphoreticular system, particularly of T-cell origin, have characteristic features. Adult T-cell leukaemia/lymphoma may present with fever, rash and hypercalcaemia as well as extensive hilar lymphadenopathy. *Mycosis fungoides* is recognizable by the cutaneous deposits, which, although initially localized as lymphomatous lesions, spread as the disease progresses to involve nodes, resulting in generalized node enlargement. *Acute lymphatic leukaemia* – especially in children – may present with diffuse lymphadenopathy. *Myeloproliferative disorders* such as chronic myeloid leukaemia, polycythaemia rubra vera and myelofibrosis are occasionally associated with lymphadenopathy.

**OTHER CONDITIONS**

Lymphadenopathy can occur in *collagenoses* (e.g. systemic lupus erythematosus and rheumatoid arthritis). *Sarcoid*, in addition to causing bilateral hilar lymphadenopathy, may result in generalized superficial lymphadenopathy. *Drugs* (e.g. phenytoin) may rarely cause lymph node hypertrophy. The unusual condition of angio-immunoblastic lymphadenopathy is often a highly malignant lymphoma, but occasionally it may be secondary to drug therapy (e.g. penicillins and sulphonamides).

*Splenomegaly* may be a feature of any cause of lymphadenopathy. In acute infections, it is usually small and soft, whereas in long-standing infections or malignancies it is often firm on palpation.

**INVESTIGATIONS**

These should be directed initially at the diagnosis of the most likely conditions, as judged by the history and examination. A full blood count will indicate the presence of a leukaemic process, or atypical lymphocytes will reflect an acute viral infection (e.g. EBV or HIV). A chest X-ray may reveal hilar lymphadenopathy or pulmonary pathology (e.g. tuberculosis). A serological search for infections (e.g. EBV and CMV) may indicate active infection if a high titre of a specific IgM is present. Although a bone marrow aspirate will reveal the presence of leukaemia, a trephine biopsy is much more reliable for diagnosing an infiltrate (e.g. carcinoma or lymphoma).

An abdominal and chest computed tomography scan is a sensitive investigation for identifying pelvic, para-aortic, coeliac, mesenteric, hilar or paratracheal node enlargement. If the initial blood tests do not provide an early diagnosis, it is essential to perform a lymph node biopsy. If the initial histology reveals reactive change only, or the presence of granulomas, it is often prudent to perform further biopsies if there is a strong clinical indication that the patient may have a malignant condition.
A macule is a flat, circumscribed patch of altered skin colour of any size. (In the past, the term was often restricted to small lesions, <2 cm across, larger lesions being termed ‘patches’). Macules must be distinguished from papules, the latter being palpable. Macules may be red (e.g. rubella), dark red (e.g. purpura), brown (e.g. freckle) or white (e.g. vitiligo) (Box M.1). (See also ERYTHEMA, p. 164.)

RED MACULES
Redness that is due to hyperaemia and that blanches with pressure is known as ‘erythema’. A widespread red macular rash occurs at some stage in many infections:

- In measles, the rash begins behind the ears and on the forehead in an ill, coughing, febrile child, with conjunctivitis and lymphadenopathy. The rash spreads over the face, neck and extremities, and comprises small pink macules that become confluent, and subsequently turn brown and desquamate. Vesicles on the buccal mucosae (Koplik’s spots) are seen just before the rash appears. The pink macules of rubella (German measles) also begin behind the ears and spread onto the face, head, neck, and trunk. Occipital lymphadenopathy is common, but fever and malaise are mild. In adults, there may be an associated arthropathy.
- In typhoid, ‘rose spots’ (0.5 cm red maculopapules) appear in crops on the abdomen, chest and back. Successive crops may come and go for 2–3 weeks.
- The macular syphilide is one of the most characteristic lesions of secondary syphilis (Fig. M.1). The eruption, named ‘syphilitic roseola’, begins as a macular mottling resembling measles, but rather more dusky and distributed over the chest, over the abdomen and – of great diagnostic importance – on the palms and soles. Eroded red patches are commonly present on the buccal mucosa. Malaise and lymphadenopathy are common, and serology will be positive. A primary chancre will often still be present. Generally, about a fortnight after its appearance, the rash begins to fade, giving place to a papular or follicular eruption on the trunk, limbs, face and neck. Vesicles are never seen, and characteristically the rash does not itch. It should be distinguished from pityriasis versicolor, in which the scaly patches can be demonstrated by scratching, and the fungus can be demonstrated microscopically; from drug reactions by their more vivid redness and the presence of itching and burning; from seborrhoeic dermatitis by its scaliness and pinkish-yellow colour; from measles by the coryza, cough and the different distribution; and from pityriasis rosea by the history of the herald patch, the distribution sparing the face and extremities and the characteristic oval lesions, each with their collarette of scaling.

**Box M.1 Macules**

<table>
<thead>
<tr>
<th>Red macules</th>
<th>White macules</th>
</tr>
</thead>
<tbody>
<tr>
<td>Exanthems</td>
<td>Post-inflammatory hyperpigmentation</td>
</tr>
<tr>
<td>- Measles</td>
<td></td>
</tr>
<tr>
<td>- Rubella</td>
<td></td>
</tr>
<tr>
<td>Typhoid</td>
<td></td>
</tr>
<tr>
<td>Drug reaction</td>
<td></td>
</tr>
<tr>
<td>Macular syphilide</td>
<td></td>
</tr>
<tr>
<td>Tuberculoid leprosy</td>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Purpuric macules</th>
<th>Xeroderma pigmentosum</th>
</tr>
</thead>
<tbody>
<tr>
<td>- Inflammatory skin diseases</td>
<td>- Peutz–Jegher’s syndrome</td>
</tr>
<tr>
<td>– Contact dermatitis</td>
<td>– Hutchinson’s freckle</td>
</tr>
<tr>
<td>– Drug reactions</td>
<td>– Flat mole (junctional naevus)</td>
</tr>
<tr>
<td>– Erythema multiforme</td>
<td>– Café-au-lait spots</td>
</tr>
<tr>
<td>– Vasculitis</td>
<td>– Neurofibromatosis</td>
</tr>
<tr>
<td>Pigmented purpuric eruptions</td>
<td>– Albright’s syndrome</td>
</tr>
<tr>
<td>– Schamberg’s purpura</td>
<td>– Mongolian spot</td>
</tr>
<tr>
<td>– Majocchi’s purpura</td>
<td>– Chloasma</td>
</tr>
<tr>
<td>Scurvy</td>
<td>– Berloque dermatitis</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Brown macules</th>
<th>Post-inflammatory hyperpigmentation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Freckles</td>
<td></td>
</tr>
<tr>
<td>– Sun-induced</td>
<td></td>
</tr>
</tbody>
</table>

**Figure M.1** Rash of secondary syphilis affecting the soles of the feet.
In the earlier stages of tuberculoid leprosy, red macular areas may appear. These are often isolated and have hyperpigmented borders, and as they expand to their maximum size of 2–10 cm in diameter, the centre of the lesions becomes depigmented and anaesthetic (Fig. M.2). In dark skins, the red macules are difficult to discern until the central depigmentation makes them obvious.

In drug reactions, itching and burning are pronounced, and lesions may later become elevated (e.g. maculopapular ampicillin rash), purpuric or even bullous.

**PURPURIC MACULES**

Purpuric macules are caused by the escape of red blood cells into the skin, and characteristically fail to blanch on pressure when a glass slide is pressed on the skin (Fig. M.3). Tiny lesions 1–5 mm are true purpura, larger extravasations are ecchymoses, and associated subcutaneous collections are haematomas. Purpura appear suddenly, are painless and change hue from red to brown, to green and yellow before fading. Where purpura is due to vasculitis, including meningococcal septicaemia, the lesions are generally palpable. Perifollicular purpuric macules may be seen in scurvy (ascorbic acid deficiency).

**BROWN MACULES**

Freckles (ephelides) are small (less than 0.5 cm), brown, roundish macules in areas of the skin exposed to the sun. They appear in childhood, especially in red-haired, fair-skinned individuals, and are associated with vulnerability to sunburn. They are particularly florid in the rare genetic syndrome xeroderma pigmentosum, a condition associated with deficient repair of sun-damaged nuclear protein. By their early teens, affected children are disfigured by atrophy, telangiectasia and multiple cutaneous malignancies, which ultimately prove fatal.

In Peutz–Jegher’s syndrome, freckles in great profusion on and around the lips (Fig. M.4a) are associated with polyposis of the small bowel, which may be complicated by recurrent intussusception and melaena. The freckles are smaller than usual and
may extend inside the mouth and onto the sides and backs of fingers. *Hutchinson's freckle* is a giant freckle that usually occurs on an elderly, sun-exposed skin. Pigmentation is variegate, and malignant pigment cell tumours often eventually arise within their boundaries. *Flat moles* (*junctional naevi*) are either present at birth or develop during the first two decades of life. They have distinct borders and are not uniformly pigmented. They undergo gradual spontaneous involution. Rarely, a more rapid involution of moles – both flat and papular – may be accompanied by a stark, chalk-white halo of depigmentation – a halo naevus (*Sutton's naevus*). This is a dramatic, but benign, condition.

*Café-au-lait spots* are well named; they are creamy brown in colour, with smooth borders, and range from 1 to 15 cm in diameter (Fig. M.4b). More than six lesions on a child's skin are said to be highly suggestive of the autosomal dominant von Recklinghausen's neurofibromatosis. The later appearance of axillary freckling is an even stronger diagnostic pointer to the later appearance of multiple pendulous cutaneous nerve sheath tumours. Large brown macules with irregular borders on a child's trunk can also be associated with bone cysts and precocious puberty in girls with *Albright's syndrome*.

The *mongolian spot* is well known to midwives as a marker of Asiatic or Negroid parentage. Large brown or slate-grey macules are seen over the sacrum at birth, and gradually fade over the first year of life; the colour is due to melanocytes in the dermis.

Macular facial pigmentation is a feature of *chloasma* (see SKIN, PIGMENTATION, p. 617).

**WHITE MACULES**

*Post-inflammatory depigmentation* may occur following inflammatory dermatoses, such as psoriasis or pityriasis rosea. Perhaps more commonly, many skin diseases fail to tan on a sunny holiday. This is particularly seen over the cheeks and outer arms of children with the eczematous condition *pityriasis alba*.

*Vitiligo* may appear at any age, presenting with symmetrical chalk-white patches of complete depigmentation around eyes, mouth, genitals and axillae. White locks of hair may occur, but ocular pigmentation is never affected. In dark-skinned people, the lesions are clearly seen (Fig. M.5), but in untanned, pale individuals the use of an ultraviolet light may be necessary to demonstrate the full extent of the disease.

*Pityriasis versicolor* is becoming a more common cause for medical consultation. The original fawny-pink, slightly scaling, round patches on the chest and back may not be noticed until, following sun exposure, partial depigmentation of affected areas occurs. The diagnosis can nearly always be made clinically by noting the slight superficial scale and looking at the margin for typical tiny annular lesions. Microscopic examination reveals the causative yeast. The most difficult differential diagnosis is post-inflammatory depigmentation following guttate psoriasis, but here the small round white macules are scattered uniformly over the body and limbs rather than concentrated in the mantle area.

Ash leaf-shaped hypopigmented macules are seen in *tuberous sclerosis*, and these may be more readily seen under ultraviolet light. *Naevus anaemicus* is present from birth and may be found at any site as a group of islands of blanched skin. The pigment mechanisms are intact, but there is permanent vasoconstriction due to a congenital neurovascular abnormality.

**MARASMUS**

Dipak Kanabar

Severe protein–energy malnutrition in children usually leads to marasmus, which is a body weight less than 60 per cent of the mean for age. Children (usually aged less than 1 year) are often withdrawn and apathetic, their facial appearance is pinched and grey, and they have a curiously senile (wizened) expression. The eyes and the fontanelle are sunken, the skin is tightly stretched over the bones of the skull, whereas on the limbs and body generally the skin is thin and inelastic through dehydration and loss of subcutaneous fat, so that it hangs in festoons on the stick-like arms and legs (skin-fold thickness and mid-arm circumferences are markedly reduced). The thorax is particularly...
wasted, and the ribs are unduly prominent. The bony fingers are often stuffed into the mouth as if to obtain nourishment. Oedema is not present. There may be a skin rash with hyperkeratosis, angular stomatitis, sparse depigmented hair, diarrhoea, hypothermia, bradycardia and hypotension, and low levels of electrolytes in the blood. These children are at great risk of intercurrent illness such as gastroenteritis, measles and tuberculosis and by incorrect feeding of carbohydrate rather than protein. Kwashiorkor, a particular manifestation of protein–energy malnutrition, occurs in some developing countries where young children’s diet is high in starch. Vitamin and mineral deficiencies also complicate the condition, making treatment far more difficult than in uncomplicated marasmus.

Conditions leading to marasmus include: starvation (from either neglect, extreme poverty, a breakdown of normal civilized behaviour, as usually occurs in modern warfare); persistent vomiting (e.g. hiatus hernia) or diarrhoea; chronic infections (of the urinary tract, tuberculosis, congenital syphilis, HIV and parasitic infestations); malabsorption – of fat in coeliac disease and cystic fibrosis, of sugar in carbohydrate intolerance, or of protein in protein-losing enteropathy; a group of conditions associated with polyuria, i.e. diabetes mellitus, diabetes insipidus, hypercalcaemia, renal acidosis and renal failure; and, rarely in children, thyrotoxicosis, Addison’s disease and malignant disease. Severe cyanotic congenital heart disease that is not corrected may be associated with a marked failure to thrive.

Starvation has a variety of causes. Because of social and political circumstances, the child may simply not receive enough calories. As a result of ignorance, feeding may be imperfect both as regards quantity and quality (see kwashiorkor, below). Because of structural imperfections (e.g. cleft lip, cleft palate) or feebleness from prematurity, the infant may be unable to suck. Oesophageal stricture and associated dysphagia may prevent adequate intake of food, and persistent vomiting will have a similar disastrous effect.

Chronic infections of the upper renal tract, and associated renal failure, for example in advanced bilateral congenital hydronephrosis, may be an aggravating factor. Congenital syphilis was once a potent cause of marasmus, although its classic features of snuffles, skin lesions, Parrot’s nodes, condylomas and enlargement of the liver and spleen are now rarely seen. Chronic infections associated with the acquired immune deficiency syndrome (HIV/AIDS) are the contemporary equivalent. Advanced miliary tuberculosis is now fortunately rare in the UK, as is neglected tuberculous disease of bone, with its associated discharging sinuses leading to secondarily infected abscess cavities.

Malabsorption is a potent cause of weight loss in children. Cystic fibrosis and coeliac disease are associated with gross steatorrhoea. Disaccharidase deficiency may result in the infant being unable to split disaccharides into absorbable monosaccharides. Lactase deficiency is the most common example, but the enzymes responsible for splitting sucrose and maltose also may be affected. Not only is the child prevented from absorbing sugar, but sugar also remains in the small intestine to aggravate the condition by causing diarrhoea by an osmotic effect. Protein-losing enteropathy due to a variety of small intestinal diseases, including Crohn’s disease, may result in protein loss into the bowel lumen.

Diabetes mellitus may have a relatively acute onset in children, with thirst, polyuria and severe loss of weight.

Kwashiorkor is a condition seen chiefly in Central Africa in children who are reared in traditional polygamous societies. It has also been widely described in other tropical and subtropical parts of the world, and even in Europe. It is a state produced by gross protein deficiency, and it is usually due to late weaning of children from the breast, and a high-starch diet. The majority of sufferers are infants under the age of 2 years who are placed on a diet mainly of cereals, with little (if any) animal protein because of shortage of suitable foods or deeply ingrained superstitions. In addition to the failure of growth, there is oedema of the legs, a distended abdomen, hyperpigmentation, and very typical depigmentation of the skin and hair, which produces a reddish hue in African babies.

MELAENA

Simon Anderson

Melaena is the term for black bowel motions resulting from haemorrhage that has occurred in the gastrointestinal tract at a high enough level for chemical alteration to take place (usually the upper gastrointestinal tract, but it could also be as far down as the caecum). Melaena may also occur after swallowing blood derived from haemoptysis or epistaxis. Melaena stools are black, tarry and with a sticky consistency, rendering them difficult to flush down the toilet. It has been shown by feeding healthy volunteer medical students with increasing aliquots of their own blood that between 50 and 80 ml of blood is sufficient to cause a melaena stool.
Black or dark stools, simulating melaena, commonly occur with iron preparations (the iron being converted to the sulphide form) and also after taking bismuth. Dark stools may also occur after eating a lot of liquorice, charcoal biscuits, black cherries, blueberries or from the excretion of large amounts of bile pigments. The characteristic thick, sticky nature of melaena stools is generally easily differentiated from other causes of black stools, but the diagnosis can be confirmed by testing the stool for the presence of blood.

Melaena is most commonly due to bleeding from the upper gastrointestinal tract (oesophagus, stomach and duodenum). If from the oesophagus or stomach, it is often associated with haematemesis, before the melaena is apparent. If melaena occurs alone from these sources, it generally indicates that the rate of bleeding is relatively slow. Melaena is as significant as haematemesis as an indication of upper gastrointestinal haemorrhage; patients with melaena should be investigated and managed as urgently as those with haematemesis. It is possible to judge the severity of a bleeding from the patient's description of the stools.

A detailed account of symptoms such as fainting, sweating and collapse, together with the general assessment of the patient's haemodynamic state, allows the clinician to assess the severity of the gastrointestinal bleed leading to the development of melaena. Upper gastrointestinal endoscopy is always needed, and a blood transfusion may be necessary.

The most common causes of melaena (perhaps more than 85 per cent of patients) include duodenal or gastric ulceration, acute gastric erosions, reflux oesophagitis or oesophageal varices (which usually also results in haematemesis). Bleeding from erosions due to non-steroidal anti-inflammatory drugs or aspirin is an increasingly common cause of melaena, especially in elderly patients.

Lesions distal to the duodenum generally give rise to dark (maroon) or bright red blood in the stools rather than melaena. However, melaena may occur from the relatively uncommon sources in the small bowel, including mesenteric ischaemia, gastrointestinal stromal tumours (GISTs) Meckel’s diverticulum, Crohn’s disease, typhoid gastroenteritis (due to ulceration of an ileal Peyer’s patch), angiodysplasia with coagulopathy, blood dyscrasias or anticoagulant therapy. Rarely, melaena may be associated with isolated small intestinal ulceration in coeliac disease or secondary to NSAIDs.

**MEMORY, DISORDERS OF**

Mark Kinirons

*(See also CONFUSION, p. 102.)*

Disorders of memory are extremely important in clinical practice. Numerous tests are used to assess memory. One of the easiest to use is the Mini Mental State Examination (MMSE), which tests orientation, registration, attention and calculation, recall, language, construction and spatial orientation (areas of higher cortical function). The MMSE has been used widely, although not in all cultures, and depends on education level. The causes of memory loss are given in Box M.2.

**FUNCTIONAL MEMORY DISTURBANCE/ PSEUDODEMENTIA**

‘Pseudodementia’ is the term used to describe non-organic causes of memory impairment. A depressed person may perform as poorly as a demented person on tests of memory simply because they are psychomotor retarded and lack the motivation to take an interest in their surroundings. Similarly, a patient who is psychotic may interpret their recall of events by delusional beliefs. Memory lapses are also common in states of fatigue. In *hysterical fugue states,*

**Box M.2 Causes of memory loss**

<table>
<thead>
<tr>
<th>Category</th>
<th>Causes</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Common</strong></td>
<td></td>
</tr>
<tr>
<td>• Head injury</td>
<td></td>
</tr>
<tr>
<td>• Cerebrovascular disease</td>
<td></td>
</tr>
<tr>
<td></td>
<td>– Multi-infarct dementia</td>
</tr>
<tr>
<td></td>
<td>– Single infarct in region of posterior cerebral artery</td>
</tr>
<tr>
<td></td>
<td>– Subarachnoid haemorrhage</td>
</tr>
<tr>
<td>• Alzheimer-type dementia (and other causes of organic confusion states; see pp. 102–4)</td>
<td></td>
</tr>
<tr>
<td>• Thiamine deficiency (Wernicke–Korsakoff syndrome) due to alcoholism, malabsorption, carcinoma of the stomach or hyperemesis gravidarum</td>
<td></td>
</tr>
<tr>
<td>• Electroconvulsive therapy</td>
<td></td>
</tr>
<tr>
<td><strong>Less common</strong></td>
<td></td>
</tr>
<tr>
<td>• Herpes simplex encephalitis</td>
<td></td>
</tr>
<tr>
<td>• Verteobasilar disease (transient global amnesia)</td>
<td></td>
</tr>
<tr>
<td>• Tumours of third ventricle or hypothalamus</td>
<td></td>
</tr>
<tr>
<td>• Depression (pseudodementia)</td>
<td></td>
</tr>
<tr>
<td>• Fugue states and psychogenic amnesias</td>
<td></td>
</tr>
<tr>
<td>• Malingering</td>
<td></td>
</tr>
<tr>
<td><strong>Rare</strong></td>
<td></td>
</tr>
<tr>
<td>• Bilateral temporal lobectomy for intractable epilepsy</td>
<td></td>
</tr>
<tr>
<td>• Intractable epilepsy</td>
<td></td>
</tr>
<tr>
<td>• Carbon monoxide poisoning</td>
<td></td>
</tr>
<tr>
<td>• Personality disorder</td>
<td></td>
</tr>
<tr>
<td></td>
<td>– Pseudologia fantastica</td>
</tr>
<tr>
<td></td>
<td>– Ganser state</td>
</tr>
</tbody>
</table>
there is a narrowing of consciousness, a move away from normal surroundings and subsequently complete amnesia for these events. The behaviour of the person may appear quite appropriate during the episode, which can last from hours to weeks. Such dissociative states may be attempts by people of certain personalities to cope with real or imagined stress. The Ganser state, first described in prisoners awaiting sentence, should also be considered a hysterical dissociative state. Two of the main features of the condition are clouding of consciousness and approximate answers (2 + 2 = 5). Subjects are subsequently amnesic for their abnormal behaviour.

**ORGANIC MEMORY IMPAIRMENT**

The most common cause of organic memory impairment is Alzheimer’s disease in 60 per cent, followed by 30 per cent of cases being due to multiple cerebrovascular accidents. In the West, tertiary syphilis is now gone, but it is an important cause in the developing world. Rare causes of memory impairment also need to be considered. Specific lesions to specific organs can cause specific types of impairment. It has long been known that damage to the temporal lobes and to structures in the limbic system, in particular the hippocampus, amygdala, fornix, mammillary bodies and dorsal medial nucleus of the thalamus, may result in profound amnesia. Bilateral focal lesions affecting these structures, for example temporal lobe damage following herpes simplex encephalitis, can cause memory impairment. Organic memory disturbance may also be a feature of diffuse cerebral disorder perhaps involving quite different brain mechanisms. In some situations, frontal lobe dysfunction may contribute to memory impairment. Any pathological process that causes bilateral damage to these diencephalic and limbic structures, such as head trauma, hypoxia, infarction in the territory of the posterior cerebral arteries or herpes simplex encephalitis, can cause memory impairment.

*Alzheimer’s disease*, first described by Alzheimer in 1907, describes a progressive decline in high critical function that eventually leads to inexorably to a mute bedridden state over 3–10 years. Initially, the patient is able to function at home, but progressive memory loss and increasing behavioural disturbance lead to an inability to carry out complex followed by simple tasks, which leads to increased care needs from family and others. Patients die from complications such as pneumonia. There is no cure. There are drugs available to help alleviate suffering in the mild to moderate stage. *Multi-infarct disease* (MID) is increasingly recognized due to the increasing incidence of cardiovascular disease and successful ageing. It occurs in patients with increased cholesterol, diabetes, hypertension or stroke disease.

The main cause of dementia is thromboembolic disease leading to occlusion of the arteries that supply the brain, which in turn leads to memory impairment. It manifests as a stepwise decline in cognitive function. There is no specific treatment. Strategies to reduce the risk factors and control conditions mentioned above are worthwhile.

*Lewy body dementia* is increasingly being recognized as a cause of dementia due to the excess deposition of Lewy bodies in the brain's subcortical structures. This leads to presenting symptoms differing from Alzheimer’s disease or MID. Typically, patients have psychomotor disturbance prominent visual hallucinations, a fluctuating level of consciousness (from day to day) and exaggerated sensitivity to major tranquillizers. The disease is incurable and progressive. Rarer forms encountered include *Wernicke–Korsakoff syndrome*, caused by thiamine deficiency in chronic alcoholism, and less frequently due to carcinoma of the stomach, hyperemesis during pregnancy, malabsorption or dietary deficiency. A similar clinical picture can present with tumours of the third ventricle and hypothalamus, carbon monoxide poisoning, tuberculous meningitis or subarachnoid haemorrhage. The acute effects of thiamine depletion include clouding of consciousness, ataxia and nystagmus (Wernicke's encephalopathy); loss of short-term memory is the principal lasting feature. Patients characteristically retain their long-term memory and can accurately recall events before the onset of the illness. They also have intact immediate memory (digit span) but will be completely unable to store this information for more than 1 or 2 seconds. With this profound loss of recent memory, patients *confabulate*, which is a way of covering up an exposed memory gap. Sometimes confabulation will become fantastic in nature as patients invent situations in their lives. Memory loss in Wernicke–Korsakoff syndrome is generally permanent when the cause is alcohol abuse.

*Transient global amnesia* describes a memory impairment that starts abruptly and improves over several hours. It generally occurs in middle-aged patients and is thought to be due to bilateral temporal lobe ischaemia caused by verteobrosasial disease. During an attack, the patient is unable to form new memories, and their behaviour may be outwardly normal. After recovery, the subject is left with a permanent amnesia for the period of the attack. *Closed-brain injury*, chiefly a result of road
traffic accidents, but also due to falls, assault and sports injuries, is a major cause of organic memory impairment. This may be due to focal or diffuse cerebral damage caused by intracranial haematoma, brain swelling, infection, subarachnoid haematoma, hydrocephalus or extracranial factors such as hypoxia and hypotension. Post-traumatic amnesia refers to the period that elapses between injury and the restoration of normal memory. The duration of post-traumatic amnesia is one indicator of the severity of head injury. Typically, patients have an impairment in immediate and recent memory following head injury, with difficulty performing digit span tests as well as tests for words and sentences. Most recovery takes place within 1 year of injury.

MENSTRUAL DISORDERS

Tony Hollingworth

‘Menstrual disorders’ is a general term to denote a wide range of presenting symptoms related to menstrual bleeding. Menorrhagia or heavy periods refers to excessive menstrual flow, or undue prolongation of the time during which it takes place. The patient is free from bleeding during the intermenstrual periods. The terms metrorrhagia (not commonly used) and irregular uterine bleeding are reserved for bleeding that occurs between the periods. The latter may be also associated with heavy periods. Pure menorrhagia is an important symptom of many well-defined conditions that do not, as a rule, give rise to irregular bleeding. Both these terms must be limited carefully to patients who menstruate, and must not be used for bleeding after the menopause. It may be more appropriate to use the term ‘dysfunctional uterine bleeding’ especially when there is no obvious underlying cause.

MENORRHAGIA

‘Heavy periods’ is a very subjective symptom, and menstrual loss consists not only of blood but also tissue and other secretions. Objectively, menorrhagia is taken to be more than 80 ml blood loss per month, which will result in iron deficiency anaemia. The diagnosis of menorrhagia may be difficult because of the absence of anaemia or other signs of severe menstrual blood loss. The diagnosis has to be accepted when the patient complains of having to use more than a dozen and a half pads per menstrual period, or when she loses clots or has episodes of flooding with blood. It is the second most common cause for hospital referrals, and up to one-third of women may consult their primary care physician about this symptom at some stage in their life.

Excessive menstrual loss in women without abnormal physical signs is believed to be endocrine in origin and is termed ‘dysfunctional menorrhagia’. Acute endometritis of gonococcal or pyogenic origin tends to cure itself owing to the shedding of the endometrium during menstruation. Tuberculous endometritis, a rare cause of infertility in the UK, is due to spread from the Fallopian tubes and is therefore associated with menorrhagia due to the tuberculous salpingo-ophoritis. If a tuberculous infection is suspected, the uterine curettings should be examined for the typical tubercles and the organism isolated by culture. Causes of menorrhagia are given in Table M.1.

Dysfunctional menorrhagia

Menorrhagia of puberty is mainly due to hypofunction of the anterior pituitary body, with a consequent failure of ovulation, and therefore no development of the corpus luteum. The ovaries contain unruptured Graafian follicles; there is increased oestrogen production, and a lack of the luteal hormone progesterone. These cases often rectify themselves in time as the pituitary gradually assumes its normal cyclical activities. These anovulatory cycles are usually painless and can be prolonged.

Menorrhagia in mature women without any obvious lesions of the generative or other systems is thought

### Table M.1 Causes of menorrhagia

<table>
<thead>
<tr>
<th>Pelvic pathology</th>
<th>Endocrine</th>
<th>Blood dyscrasias</th>
<th>Iatrogenic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fibroids</td>
<td>Anovulatory – usually at extremes of reproductive age</td>
<td>Thrombocytopenia</td>
<td>Use of intrauterine contraceptive device (IUCD; especially copper types)</td>
</tr>
<tr>
<td>Pelvic inflammatory disease (PID)</td>
<td>Thyroid disease, especially myxoedema</td>
<td>Von Willebrand's disease</td>
<td>Progesterone-only</td>
</tr>
<tr>
<td>Endometriosis and adenomyosis</td>
<td>Cushing's disease</td>
<td>Christmas disease</td>
<td>Contraception</td>
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<tr>
<td>Tuberculosis of the pelvis</td>
<td>Anticoagulation</td>
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<td></td>
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<tr>
<td>Malignancy</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pregnancy related</td>
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</tbody>
</table>
to be due to an imbalance between the secretion by the ovary of oestrogen and progesterone, with an increase in oestrogen and a complete lack of or a deficiency of progesterone. When there is a complete absence of progesterone in the second half of the menstrual cycle, the cycle is referred to as anovular, drawing attention to the failure of ovulation and formation of a corpus luteum. Sometimes the ovaries become cystic, and the endometrium undergoes polypoidal thickening with a characteristic microscopic appearance known as ‘Swiss cheese’ endometrium or cystic glandular hyperplasia. This condition has been known as metropathia haemorrhagica in the past. Bouts of amenorrhoea of some weeks are followed by prolonged irregular bleeding, a symptom complex that does not properly come under the heading ‘menorrhagia’.

Menorrhagia in relation to the menopause and in the years preceding it is the result of an increasing failure of ovarian function and consequent upset in the balance between the secretion of oestrogen and progesterone. {

Polymenorrhoea is the name given to a form of irregular and excessive menstruation in which the cycle is shortened from the usual 28 days to 21 days or even less. This is due to a disturbed balance of internal secretions, causing ovulation to occur too early in the cycle; in some cases, two corpora lutea have been found at the same stage of development. In many instances, fibroids are present.

Pelvic pathology
Menorrhagia can be associated with fibroids (benign leiomyomas), adenomyosis, pelvic infection, endometrial polyps, endometriosis and the presence of an intrauterine contraceptive device (IUCD). Of all the causes of pure menorrhagia, leiomyoma (fibroids) of the uterus stands out as the only important growth associated with this symptom, and a simple bimanual examination, as a rule, suffices to show that such a tumour exists. The size and shape of the uterus is dependent on the number and size of the fibroids, as there may be more than one myoma in the uterus, causing its shape to be exceedingly irregular; the uterus feels firm, and in most cases it is mobile. The only difficulty in diagnosis, as a rule, lies in distinguishing a leiomyoma of the uterus from an ovarian cyst. Ultrasound scanning is helpful in the diagnosis of fibroids because it is possible by this method to determine the origin and nature of a pelvic swelling. Fibroids may be submucous, intramural, subserosal or pedunculated. Submucous fibroids distortion of the uterine cavity with an increase in the endometrial surface area from which menstruation occurs leading to menorrhagia (Fig. M.6).

Chronic salpingo-oophoritis (in the form of a pyosalpinx, a hydrosalpinx, a tubo-ovarian abscess or chronic interstitial salpingitis) and ovarian endometriosis both give rise to menorrhagia due to pelvic congestion, but dysmenorrhoea, pelvic pain, dyspareunia and backache are usually more
prominent symptoms. In either case, a firm tender swelling in the pouch of Douglas can be felt on bimanual palpation. It is unusual to get menorrhagia in these conditions without bleeding at other times of the cycle.

Adenomyosis is a condition that can present with menorrhagia and pain at the time of menstruation; on bimanual examination the uterus may be tender. The diagnosis can only be confirmed histologically as endometrial tissue is found in the myometrium. The condition is more common in parous women, and in essence there has been bleeding in the myometrium that gives rise to the pain and tenderness (Fig. M.7).

Anticoagulation in women who are on long-term anticoagulants for prosthetic heart valves, previous pulmonary embolism or, in some cases, antiphospholipid syndrome may develop significant period problems depending on the value of the International Normalized Ratio (INR) that is required therapeutically.

Severe menorrhagia may complicate thrombocytopenia, but as soon as the underlying cause is treated, the period loss becomes normal.

**Medical disorders**

The function of the thyroid and suprarenal glands can influence the menstrual loss, although the mechanism is unknown. Menorrhagia tends to be more common in hypothyroidism than thyrotoxicosis, and it is not uncommon with Cushing’s disease.

**Irregular uterine bleeding**

**IRREGULAR UTERINE BLEEDING (METRORRHAGIA)**

Metrorrhagia means loss of blood vaginally between the menstrual periods, and the term should be applied strictly only to irregular haemorrhages during the reproductive age range, i.e. from puberty to menopause. It may be used for losses of actual blood or for blood-stained discharges in which mucus is mixed with blood. There has been a tendency of late to refer only to ‘menorrhagia’ (see above) or use the term ‘dysfunctional uterine bleeding’. ‘Metrorrhagia’ as a term is no longer commonly used in clinical practice. However, it is a term that includes all types of irregular vaginal bleeding, whether it occurs during menstrual life, before puberty, after the menopause or during pregnancy. For the purposes of this discussion, irregular vaginal bleeding will be considered under three headings:

- Irregular bleeding during menstrual life
- Irregular bleeding before puberty and after the menopause
- Irregular bleeding during pregnancy

It is important to emphasize that, if a woman has had regular periods and then starts to get irregular bleeding for no apparent reason, one must exclude pregnancy and where that pregnancy is located, i.e. whether she could have an ectopic pregnancy, as this is still a major cause of maternal death in the UK.

**Irregular bleeding during menstrual life**

Causes of irregular bleeding are given in Box M.3.

**Malignancy**

Carcinoma of the cervix Cervical cancer is an uncommon disease with an incidence that is reducing in the developed world as a result of cervical...
screening programmes. It is estimated that a general practitioner in the UK will see one case of cervical cancer every 7–9 years. The cervix is replaced with a friable mass, which causes irregular bleeding as well as postcoital bleeding. A number of women will be diagnosed in the colposcopy clinic having had an abnormal cervical smear test result, although the screening programme is designed to pick up premalignant disease, which is asymptomatic. **Carcinoma of the uterine body** Endometrial carcinoma does occur during the reproductive age group, but it is much more common in postmenopausal women. It is now the most common genital tract malignant tumour in England and Wales, and it can present with irregular bleeding. The risk factors for this condition include obesity (raised body mass index (BMI)), nulliparity and a history of polycystic ovary syndrome (PCOS). It is unusual to diagnose this condition before the age of 40 years, hence the Royal College of Obstetricians and Gynaecologists’ recommendations suggesting that women with menstrual irregularities before the age of 45 years should receive treatment for 3 months. If the irregularity persists and especially if the woman is anaemic, a hysteroscopy and endometrial sampling should be undertaken; if the woman is over the age of 40 years, this would be a first-line investigation. **Sarcoma of the uterus** This is a very uncommon tumour and occurs in fibroids. It may present with irregular bleeding, but many of these women will be postmenopausal and present with a rapidly expanding pelvic mass. The risk of a fibroid becoming malignant is estimated at about 1 in 1000. This tumour may occur in an existing fibroid or appear de novo. The difficulty with this condition is the highly aggressive nature of the disease, which does not, as a rule, respond well to radiotherapy or chemotherapy.

**Chorionic carcinoma** This condition is fortunately rare, and follows hydatidiform mole in about 5 per cent of recorded cases. It always follows pregnancy, never having been seen in a uterus where pregnancy could be excluded, although the pregnancy may have occurred some years earlier. It is associated with profuse bleeding and the rapid development of a fetid discharge due to decomposition of blood and necrotic tissues in utero. Secondary deposits of chorionic carcinoma can appear as small plum-coloured, ulcerating nodules in the vagina, and secondaries in the lungs cause haemoptysis. The patient rapidly becomes ill, with pyrexia and profound anaemia. A raised level of chorionic gonadotrophin is found in the urine. The diagnosis depends upon the finding of masses of trophoblastic cells in uterine curettings without any evidence of villous formation.

**Other malignancies** Carcinoma of the Fallopian tube is a rare tumour and tends to present in postmenopausal women, although it may present with irregular bleeding. Ovarian cancer is unlikely to cause bleeding unless it has invaded the uterus. Clear-cell carcinoma of the vagina is also rare and has been reported in teenage girls exposed to diethylstilboestrol in utero.

**Benign lesions**

**Fibroids** Leiomyomas may cause a mixture of menorrhagia and metrorrhagia. If irregular bleeding occurs, the fibroids are usually submucous. They may be in the process of extrusion when they may become infected, and sloughing occurs. The reason for this is that, in these conditions, the tumours are partly strangulated by uterine contractions and consequently congested with venous blood. This results in bleeding that is unpredictable in timing and amount.

**Polyps** Polyps (Fig. M.8) can occur within the cervix and endometrium. Cervical polyps are usually asymptomatic and identified at the time of taking a routine cervical smear test, but if the tip becomes inflamed, it can give rise to vaginal bleeding or postcoital bleeding. Polyps within the endometrial cavity, whether fibroid or mucous, are common causes of intermenstrual bleeding and are usually quite definitive growths. The mucous polyp is soft, strawberry-red in colour and pedunculated, and it contains cystic spaces filled with glairy mucus. It rarely gives rise to a malignant growth, although it can become hyperplastic. A fibroid polyp is hard and shows the glistening, whorled appearance so well known in fibromyomas on section. These growths are liable to infection and sloughing and are then apt to be mistaken for carcinoma or sarcoma macroscopically.
Menstrual Disorders

Inflammatory lesions
Endometriosis This condition is defined as the finding of tissue outside the uterus that is histologically similar to that of endometrium, and it is not strictly an inflammatory lesion. However, it is one of the most common benign gynaecological conditions, and it may present with a myriad of symptoms, which include irregular vaginal bleeding, pain and dyspareunia. It remains a condition that can take a long time to diagnose, and many cases require a laparoscopy for confirmation. In some cases an MRI may be needed for identifying rectovaginal deposits.

Tuberculosis Tuberculosis may affect the genital tract and give rise to irregular bleeding and infertility. It is an uncommon problem in the UK, but is much more common in the developing world. Histology of the endometrial curettings may give the diagnosis.

Dysfunctional uterine bleeding
Although modern techniques of ovarian steroid estimation in serum allow serial measurements to be made of hormone levels throughout the menstrual cycle, no clear pattern has emerged to explain the mechanism underlying dysfunctional uterine bleeding. Dysfunctional bleeding may occur at any age between puberty and the menopause, but 50 per cent of cases occur between the ages of 40 and 50, about 10 per cent at puberty, and the remainder between these ages. The bleeding is more commonly menorrhagia, although the interval between the episodes of bleeding may be shortened. This is particularly the case when this type of bleeding occurs at the time of puberty and the menopause. The bleeding may be profuse or only slightly in excess of normal. In other cases, intermenstrual bleeding occurs, continuing for days or weeks. It is usually preceded by amenorrhoea for some weeks. In a large proportion of these cases, ovulation fails to occur, i.e. the cycle is anovulatory. The histology of any curettings may prove to be essentially normal.

Endometrial hyperplasia can occur as a result of excess oestrogen production and should be considered a potentially premalignant condition. It is likely to occur in women who are obese, due to excess aromatization of androgens to oestrogens within the adipose tissue, and it can also occur in women with unruptured follicles, as in PCOS. This may lead to irregular shedding of the endometrium. Endometrial hyperplasia is classified as:

• Simple hyperplasia without atypia
• Complex hyperplasia without atypia
• Simple atypical hyperplasia
• Complex hyperplasia with atypia, which has a greater than 20 per cent risk of malignancy

These conditions are histological diagnoses, and how the woman is treated will depend on her age and thoughts about fertility.

Contraceptive use
There are three main areas that can give rise to menstrual problems:

• Progesterone contraception, whether as the progesterone-only pill, the Mirena IUCD or Depo-Provera injection or implants, will usually result in the woman being amenorrhoeic. There are a number of women who develop irregular bleeding that may be completely unpredictable.

• The copper-containing IUCDs can give rise to menorrhagia, and in some cases the low-grade inflammatory response of the endometrium to the coil can result in irregular shedding. The treatment is to remove the coil in the first instance.

• The combined oral contraceptive pill usually gives good cycle control, but breakthrough bleeding can occur due to gastrointestinal upset, absorption and metabolic problems due to other medications such as antibiotics and antiepileptic drugs, diet affecting the enterohepatic circulation, and a dosage too low for the individual.

Bleeding associated with ovulation
It is not uncommon for women to bleed very slightly about midway between the periods at the time
Bleeding due to granulosa cell tumour

When irregular bleeding occurs in the presence of an ovarian swelling, the possibility of a granulosa cell tumour arises. Removal of the tumour and histology will reveal its nature.

Irregular bleeding before puberty and after the menopause

The causes of irregular bleeding before puberty and after the menopause are given in Box M.4. The vaginal bleeding that occurs occasionally in newborn infants is usually due to a high concentration of oestrogen in the fetal circulation. It is usually trivial, but a fatal case has been reported. Bleeding later in childhood may be due to sexual precocity, when secondary sexual characteristics will be in evidence, or due to a new growth such as an embryonal rhabdomyosarcoma (sarcoma botryoides). Vaginoscopy under anaesthesia (and biopsy if a lesion is found) is essential.

After the menopause, the differentiation of malignant growths, polyps and senile endometritis can be established only by uterine curettage. Carcinoma of the body of the uterus (endometrial adenocarcinoma) is the most common malignant growth after the menopause. In any doubtful case, routine hysteroscopy with dilatation and curettage/endometrial biopsy of the uterus must never be omitted. Senile (atrophic) vaginitis must not be overlooked as a possible cause; the vaginal walls at the fornices become inflamed and may bleed if the surfaces rub together, the surfaces may be partly adherent, and the separation brought about by the examining finger may cause bleeding. Pyometra, or distension of the uterus with pus, may cause haemorrhage, with a foul discharge; although it is very often associated with malignancy, it may be only the result of infection or cervical stenosis. The only growth of the ovary that produces uterine haemorrhage is the granulosa cell tumour. This may occur at almost any age. (See PELVIS, SWELLING IN, p. 498.)

In women with postmenopausal bleeding, ultrasound scanning to measure the endometrial thickness may be a useful way to triage these patients. If the endometrial thickness is 5 mm or less, no further action is needed unless the bleeding continues or there are repeated episodes. Otherwise, hysteroscopy and endometrial sampling should be performed.

Irregular bleeding during pregnancy

In relation to a recent pregnancy, haemorrhage may result from simple subinvolution, from retained products of conception or from choriocarcinoma. The differentiation of these conditions can be established only by exploration of the uterine cavity, with, if necessary, the assistance of histology. Such conditions may be termed ‘secondary postpartum haemorrhage’ in cases occurring within a few days of delivery.

Haemorrhage from the pregnant uterus almost always means separation of the placenta or of the embryo from its attachments, but malignant growth of the cervix, ectropions and polyps may need to be considered and so visualization of the cervix is important in these women. Haemorrhage from a pregnant uterus is never due to malignant growth of the body of the organ, because pregnancy is impossible with this lesion. There are, however, two great difficulties in connection with pregnancy haemorrhages; these are to differentiate:

- The uterine haemorrhage that occurs with extraterine (ectopic) gestation from that due to threatened miscarriage
- The bleeding of placenta praevia from that due to the separation of a normally situated placenta

In the first case, an ectopic pregnancy usually presents within the first 6–8 weeks of the last normal period. The external haemorrhage occurs when the extraterine gestation is separated from its tubal or other attachments, or is converted into a tubal mole, when it becomes extruded from the fimbriated extremity of the tube, or when the tube ruptures. These events can cause acute pain in the lower part of the abdomen, faintness and possibly collapse from internal haemorrhage. On examination, the uterus may not appear enlarged, but there may be fullness and marked tenderness in the adnexa. In the case of an ectopic gestation, the abdominal pain can be severe and is

<table>
<thead>
<tr>
<th>Box M.4 Causes of irregular bleeding</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Before puberty and after menopause</strong></td>
</tr>
<tr>
<td>Uterine bleeding in the newborn</td>
</tr>
<tr>
<td>Malignant growth of the uterus</td>
</tr>
<tr>
<td>Polyps</td>
</tr>
<tr>
<td>Senile endometritis</td>
</tr>
<tr>
<td>Senile atrophic vaginitis</td>
</tr>
<tr>
<td>Pyometra</td>
</tr>
<tr>
<td>Granulosa cell tumour of the ovary</td>
</tr>
<tr>
<td>Oestrogen withdrawal bleeding</td>
</tr>
<tr>
<td><strong>During pregnancy</strong></td>
</tr>
<tr>
<td>Threatened, inevitable or incomplete abortion</td>
</tr>
<tr>
<td>Carneous mole</td>
</tr>
<tr>
<td>Hydatidiform mole</td>
</tr>
<tr>
<td>Antepartum haemorrhage</td>
</tr>
<tr>
<td>Secondary postpartum haemorrhage</td>
</tr>
<tr>
<td>Subinvolution</td>
</tr>
<tr>
<td>Choriocarcinoma</td>
</tr>
<tr>
<td>Extraterine gestation</td>
</tr>
<tr>
<td>Malignant growth of cervix or vagina</td>
</tr>
<tr>
<td>Ectropion</td>
</tr>
<tr>
<td>Polyps</td>
</tr>
</tbody>
</table>
often referred to the shoulder. It is much more severe than that experienced in an intrauterine miscarriage, and it almost always precedes the onset of vaginal bleeding; on the other hand, the vaginal blood loss in an inevitable miscarriage is much more than that in ectopic gestation, which is usually scanty. However, these signs are not invariable.

Haemorrhage due to threatened miscarriage cannot be diagnosed unless the presence of an intrauterine pregnancy can be established, and ultrasound scanning has altered the management of these patients. The presence of a live fetus can be demonstrated by the beating of the heart with the aid of real-time ultrasound, and means that the outlook for continuation of the pregnancy is good. The presence of an empty uterus and a positive pregnancy test with an adnexal mass and fluid in the pouch of Douglas suggests an ectopic pregnancy until proved otherwise. In the case of a positive pregnancy test using methods to detect low levels of beta-human chorionic gonadotrophin (hCG) and a negative ultrasound scan for an intrauterine pregnancy, laparoscopy is advisable to exclude the diagnosis. The diagnosis of inevitable miscarriage depends upon finding some part of the uterine contents presenting through the dilating cervix. Incomplete miscarriage is diagnosed by the continuation of bleeding or seeing that not all the products of conception have been passed. Retained products are then confirmed on ultrasound scanning.

The management of this condition has changed over the past few years, with more and more women undergoing a natural miscarriage or medical management with no curettage. Curettage is reserved for women with excessive or prolonged bleeding. A hydatidiform mole (Fig. M.9) should be suspected when a rapid increase in size of the uterus occurs during the early months of pregnancy, associated with uterine bleeding and excessive symptoms of pregnancy. Most women have a uterus that is larger than would be expected for the dates, but in one-third it is smaller. Sometimes vesicles are passed and the diagnosis is clear. If not, the finding of a high level of hCG in the blood or urine and the characteristic ‘snowstorm’ appearance on ultrasound scanning makes the diagnosis certain. In a normal pregnancy, up to 100,000 IU of hCG is passed in the urine daily; five times this amount is passed in the presence of a hydatidiform mole. These patients should be registered with one of the three registration centres in the UK (Charing Cross, Sheffield and Dundee). The centres will monitor the patient for the requisite amount of time, which is dependent on how quickly the hCG level returns to normal.

Bleeding due to placenta praevia generally does not occur until 30 weeks of pregnancy. Antepartum haemorrhage is likely to be due to placenta praevia if the fetal presenting part is high above the pelvic brim, or there is a malpresentation such as a breech or a transverse lie. If the fetal head is engaged in the pelvis, antepartum haemorrhage cannot be due to placenta praevia and must therefore be due to accidental haemorrhage, provided that incidental causes such as carcinoma or a polyp on the cervix can be excluded. The diagnosis of antepartum haemorrhage has been made much easier with the aid of ultrasound scanning because the placental echo can be seen clearly, and the relation of the edge of the placenta to the internal os can be accurately determined. It is not uncommon for a low-lying placenta found in the mid-trimester to be seen on serial scanning to move wholly into the upper uterine segment as term approaches, and the lower segment develops beneath it.

In all these women, consideration needs to be given to the administration of anti-D if they are Rhesus negative.

INVESTIGATIONS
The investigations for menstrual disorders may include:

- A full blood count
- Thyroid function tests, reserved for those women who have other clinical signs and symptoms suggestive of thyroid disease
- A pregnancy test where appropriate
- An ultrasound scan of the pelvis where indicated
- Hysteroscopy and endometrial biopsy
- Diagnostic laparoscopy where indicated by the history and clinical signs
- MRI is usually the imaging of choice after ultrasound when there needs to be further delineation of a pelvic swelling
MENSTRUAL PERIODS, ABSENT

Tony Hollingworth

Amenorrhoea can be defined as the absence of menstruation, and it can be either temporary or permanent. It may occur as a normal physiological event before puberty, as a result of pregnancy and subsequent lactation, or at the onset of the menopause. It may be a symptom of a non-physiological problem that may be systemic or gynaecological in origin (Box M.5).

PRIMARY AMENORRHOEA

This is the failure to menstruate by the age of 16 years when the girl has developed normal secondary sexual characteristics, or the failure to menstruate at the age of 14 years in the absence of any secondary sexual characteristics. This definition aids the diagnostic differentiation of causes, which include reproductive tract anomalies, gonadal quiescence and gonadal failure. Primary amenorrhoea may result from congenital abnormalities in the development of the ovaries, genital tract or external genitalia, or from a disturbance of the normal endocrinological events at the time of puberty. Some of these structural abnormalities may lead to crypto-menorrhoea, where menstruation is taking place but the menstrual flow is unable to escape due to some closure of part of the genital tract.

Most causes of secondary amenorrhoea can also cause amenorrhoea if the problem occurs before puberty. Delay in the onset of puberty is often constitutional. It is important to exclude the possibility of primary ovarian failure or dysfunction of the hypothalamo–pituitary axis. As a general rule, 40 per cent of cases of primary amenorrhoea are due to endocrine disorders, and the remaining 60 per cent are due to developmental abnormalities. It is important to emphasize that the prevalence of primary amenorrhoea is around 0.3 per cent in the general female population.

SECONDARY AMENORRHOEA

The definition of secondary amenorrhoea has usually been taken to be the cessation of menstruation for 6 consecutive months in a woman who has had regular periods, although it has recently been suggested that cessation of the periods for 3–4 months may be considered pathological and warrant investigation. Secondary amenorrhoea has a prevalence of 3–4 per cent of the female population.

Irrespective of the type of amenorrhoea, a thorough history and examination should be undertaken. This should include contraceptive history, exercise levels, ...
any significant change in body mass index (BMI), eating habits and any recent stressful events, as well as any increase in body hair, any discharge from the nipple and any episodes of hot sweats (vasomotor symptoms). Examination needs to include the stature and body form of the individual; the height and weight should be measured and converted into a BMI \[\text{BMI} = \frac{\text{weight in kg}}{(\text{height in metres})^2}\].

Inspection should concentrate on the presence or absence of secondary sexual characteristics and the appearance of the external genitalia. It is essential that this be undertaken before requesting any investigations. Most cases of secondary amenorrhoea would by definition exclude congenital anomalies unless the individual had been using the oral contraceptive pill, which would induce a withdrawal bleed each month. Vaginal examination may be inappropriate in someone under the age of 16 years or someone who has not been sexually active. Abdominal ultrasound scanning is very useful to define the anatomy. It is always important to exclude pregnancy. Serum investigations should include a human chorionic gonadotrophin level, prolactin, gonadotrophins (follicle-stimulating hormone (FSH) and luteinizing hormone (LH)) and thyroid function tests.

Raised serum prolactin levels to over 1500 IU/l may indicate the need for a computed tomography (CT) or magnetic resonance imaging (MRI) scan of the pituitary fossa to exclude a hypothalamic tumour. Serum FSH levels of over 40 IU/l usually suggest irreversible ovarian failure. Raised serum FSH and LH levels usually suggest ovarian failure, but raised LH levels alone may indicate polycystic ovary syndrome (PCOS), which can be confirmed by an ultrasound scan of the ovaries. Amenorrhoea in PCOS is secondary to acyclical ovarian activity and continuous oestrogen production. Abnormally low serum levels of FSH and LH suggest failure at the level of the hypothalamus and pituitary, giving rise to hypogonadotrophic hypogonadism. Kallman’s syndrome is associated with hypogonadotrophic hypogonadism, and these patients have hyposmia and/or colour blindness. Hormonal patterns in amenorrhoea with their associated diagnoses are shown in Table M.2.

<table>
<thead>
<tr>
<th>Condition</th>
<th>Serum biochemistry</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ovarian failure</td>
<td>Raised FSH and LH</td>
</tr>
<tr>
<td>Polycystic ovary syndrome</td>
<td>Raised LH, raised free androgen index</td>
</tr>
<tr>
<td>Hypogonadotrophic/hypogonadism</td>
<td>Low FSH and LH</td>
</tr>
</tbody>
</table>

The free androgen index is the relationship or ratio of the total testosterone concentration (slightly raised) to the sex hormone-binding globulin (SHBG) concentration multiplied by a constant usually 100. It is often raised in severe acne, male androgenic alopecia and hirsutism, as well as PCOS, for which it can be a sensitive and specific indicator if elevated in the early follicular phase. In PCOS, the SHBG is lowered and the free testosterone is raised.

Chromosomal abnormalities (e.g. Turner’s syndrome – 45,XO) can be diagnosed by karyotyping. Autoantibody screens should be undertaken in women with a premature menopause. Premature menopause can be associated with an increased risk of heart disease, and consequently it may be useful to check serum cholesterol levels in these patients. Women with PCOS and prolonged amenorrhoea have an increased risk of endometrial hyperplasia and carcinoma, so endometrial sampling may be useful if any abnormal bleeding occurs.

CAUSES OF PRIMARY AMENORRHOEA

Chromosomal

In Turner’s syndrome (gonadal dysgenesis), there is also dwarfism, web neck, cubitus valgus and an XO sex chromosome pattern (Fig. M.10). This is the most common form of gonadal dysgenesis, and these...
women may develop spontaneous menstruation, although premature ovarian failure is common. The gonadotrophin levels may be raised, and women may require hormone replacement therapy (HRT). Although spontaneous conceptions have been reported, some form of assisted conception is likely to be required for women who desire a pregnancy. In testicular feminization (which is in reality androgen insensitivity), the form is female with well-developed breasts but absent or sparse pubic and axillary hair, and the gonad, which may be found in the groin or in the abdomen, is a testicle. The gonadal tissue should be removed because of the increased risk of malignancy. In ovarian dysgenesis (Fig. M.11), there are streak ovaries, an infantile uterus and absent secondary sexual characteristics. In these cases, a buccal smear for sex chromatin and a chromosome analysis on a sample of peripheral blood are indicated. In ovarian dysgenesis, there is a chromatin-negative smear but only 46 chromosomes; a single X chromosome (XO); in testicular feminization, the smear is also chromatin-negative but there are 46 chromosomes (XY). Gonadal biopsy is also helpful in diagnosis.

**Müllerian duct abnormalities**

The Wolffian ducts regress in the embryo after the sixth week if there is no Y chromosome present. The Müllerian ducts will develop into the tubes and uterus, and fuse caudally with the urogenital sinus to produce the vagina (Fig. M.12). Abnormalities may occur in the process of fusion, and these may be medial or vertical, and give rise to primary amenorrhoea. Complete or partial Müllerian agenesis may occur. In these cases, the genotype is 46XX with normal secondary sexual characteristics and normal ovarian tissue, but the vagina is short and may require surgery. There may also be associated urinary tract abnormalities.

The most common form of abnormality is that of an imperforate hymen, which leads to primary amenorrhoea or cryptomenorrhoea (hidden menses). The secondary sexual characteristics are normal, and the individual may complain of cyclical lower abdominal pain and abdominal distension. It is not unusual for these cases to present with retention of urine and on inspection to have a bulging hymen (Fig. M.13). A cruciate incision releases the menses, and that is all that is necessary.

**Figure M.11** Characteristics of ovarian dysgenesis.
CAUSES OF SECONDARY AMENORRHOEA

Genital tract abnormalities

There is potential for scarring anywhere within the genital outflow tract. Asherman's syndrome is a condition in which intrauterine adhesions develop that prevent normal endometrial growth. It is an uncommon condition, and it usually occurs following vigorous curettage at the time of an evacuation of the uterus or suction termination of pregnancy.

Cervical stenosis can cause cryptomenorrhoea with the development of a haematometra. It may occur due to repeated treatment of the cervix for precancerous lesions. Radiotherapy may have an effect on the cervix and uterus if used for advanced cancer of the cervix, and this may possibly cause vaginal stenosis. In these cases, the amenorrhoea is more likely to be related to the radiotherapy effect on the ovaries than to outflow obstruction.

Systemic disorders

Chronic disease may cause menstrual disorders as a consequence of the general disease state, weight loss or effects on the hypothalamic–pituitary axis. Certain disorders will affect gonadal function directly.

Figure M.12 Development of the genital tract.

Figure M.13 Imperforate hymen.
Chronic renal disease will act by increasing the serum LH level, and also increasing the prolactin levels, possibly due to reduced renal clearance. Other causes include systemic conditions in the form of tuberculosis or sarcoid.

**Weight-related amenorrhoea**
Body weight/BMI can have a significant effect on the regulation and release of serum gonadotrophins. Menstruation will not occur regularly if the BMI falls below 18, and it is estimated that 22 per cent of female body weight should be fat to ensure ovulatory cycles. Fat in the form of adipose tissue is a source of oestrogen by the aromatization of androgens to oestrogen. This ensures the appropriate feedback mechanism of the hypothalamic–pituitary–ovarian (HPO) axis. The weight loss may be due to illness, exercise or dieting. Potential sequelae of a low BMI also include long-term effects on bone mineralization.

Stress in itself is unlikely to give rise to amenorrhoea lasting longer than 2 months unless it is associated with debilitation. Exercise, particularly in endurance events, is a common cause of amenorrhoea, and this is also usually related to the BMI and the body fat content, as described above.

**Hypothalamic causes**
These causes are uncommon and include craniopharyngioma, gliomas and dermoid cysts. The mechanism of action may be to destroy local tissue or disrupt dopamine production, resulting in hyperprolactinaemia. Treatment is usually surgical or radiotherapy. HRT may be necessary, together with other hormonal supplementation, depending on the extent of the local damage in the pituitary. Head injury or irradiation may have a similar effect.

**Pituitary causes**
The most common pituitary cause of amenorrhoea is hyperprolactinaemia; this may be physiological due to lactation, iatrogenic or pathological. A non-functioning tumour or pituitary adenoma may affect dopamine secretion levels, as may prothiazine and metoclopramide. The consequence is a rise in the serum prolactin level. Galactorrhoea may occur in up to a third of patients, and very occasionally there may be visual field impairment.

Unless the serum prolactin is markedly raised, it is unlikely to show any effect on the sella turcica on a lateral skull X-ray. CT or MRI scanning may be a more appropriate investigation. Treatment involves the use of a dopamine antagonist, usually bromocriptine or a related drug. This should be discontinued if the patient becomes pregnant, as a quarter of the adenomas will increase in size during pregnancy.

Profound hypotension following delivery can cause Sheehan’s syndrome, which affects the pituitary to cause necrosis, as the blood supply is an end-artery with no collateral supply to protect it in these circumstances. Appropriate induction agents will be needed to induce ovulation.

Treatment needs to be given to correct the amenorrhoea and oestrogen deficiency, improve libido and effect tumour shrinkage in cases with hyperprolactinaemia. It is safe to use the combined oral contraceptive pill in these women if they require contraception.

**Ovarian causes**
Premature ovarian failure may occur, defined as the cessation of periods before the age of 40 years. This may be due to chromosomal abnormalities, which have already been discussed, and also chromosomal mosaicsisms. The most common causes include autoimmune disease, as well as infection, previous surgery, chemotherapy and radiotherapy.

Tumours are an unusual cause of amenorrhoea, but arrhenoblastomas can cause virilism as well as amenorrhoea, atrophy of the breasts and hirsutism.

**Iatrogenic causes**
The obvious ones include radiotherapy and chemotherapy for malignant disease. Others that may need to be considered are forms of contraception, including Depo-Provera, the progesterone-only pill and the Mirena coil, post-pill amenorrhoea and gonadotrophin-releasing hormone analogues. Box M.6 lists drugs that have also been associated with secondary amenorrhoea.
MENSTRUAL PERIODS, INFREQUENT (OLIGOMENORRHOEA)

Tony Hollingworth

‘Oligomenorrhoea’ is a term that defines menstrual periods that occur repeatedly at intervals between 6 weeks and 6 months. It is an arbitrary definition and may be misleading. It is considered that the normal menstrual cycle has an upper limit of 35 days. The proliferative phase (i.e. the time during which the follicle develops) is variable; the secretory phase (the time from ovulation to menstruation) is usually constant at 14 days. Cycles of 6 weeks’ duration seem to show no difference from normal-length cycles from the point of follicular and hormone development.

There is a range of conditions that can cause oligomenorrhoea, some of which can also cause amenorrhoea (see p. 409). Some common causes are:

• Polycystic ovarian syndrome accounts for about 90 per cent of cases of oligomenorrhoea, compared with only 33 per cent of cases of amenorrhoea. The periods are usually light, and the condition is often associated with anovulation.
• A prolonged proliferative phase is associated with ovulatory oligomenorrhoea. It often occurs in adolescent girls or at the time of menarche, and in older women in the perimenopausal phase.
• Prolonged corpus luteum activity may also lead to oligomenorrhoea and a prolonged cycle, but this is usually associated with prolonged menstruation.
• Exercise, weight loss and stress.

Clinically, oligomenorrhoea should be considered in the same way as amenorrhoea for investigations and further management.

MENSTRUAL PERIODS, PAINFUL (DYSMENORRHOEA)

Tony Hollingworth

The term ‘dysmenorrhoea’ comes from the Greek meaning difficult monthly flow; however, it is taken to mean painful menstruation. It is a symptom complex that includes cramping lower abdominal pain radiating to the back and legs, associated with some gastrointestinal upset, malaise and headaches. The problem can be divided into:

• Primary dysmenorrhoea, when the periods are painful and no organic or psychological cause can be found. It usually occurs at the beginning of reproductive life when the girl starts ovulating. The pain starts with the onset of menstruation and is generally associated with ovulatory cycles. There is an abnormally high production of endometrial prostaglandins, which causes excessive uterine contractions. Examination findings are usually normal, and further investigation may only be necessary if treatment fails to alleviate the symptoms. The options for treatment include the combined oral contraceptive pill to inhibit ovulation, or non-steroidal anti-inflammatory agents, which act as prostaglandin synthetase inhibitors to decrease the concentration of local prostaglandins and thereby reduce pain and also menstrual loss.
• Secondary dysmenorrhoea occurs when the woman experiences painful periods for which an organic or psychosexual cause can be found. The differential diagnosis includes:
  – Pelvic inflammatory disease
  – Endometriosis or adenomyosis
  – Fibroids
  – An intrauterine contraceptive device
  – Cervical stenosis following treatment for precancer
  – An ovarian tumour
  – Previous pelvic or abdominal surgery
  – A previous history of sexual abuse or other psychological problems

A detailed history is important, and this may take time if there is a psychosexual element. Pelvic examination should be performed, and swabs taken if indicated. A tender uterus will usually indicate the possibility of adenomyosis (see Fig. M.7). Restricted mobility or a fixed retroverted uterus may suggest the presence of adhesions secondary to endometriosis, pelvic inflammatory disease or previous surgery. A previous history of cone biopsy or other excision procedures for cervical intraepithelial neoplasia might suggest the possibility of cervical stenosis, and may require dilatation of the cervix.

Investigations will depend on the history and may include:

• Triple swabs if infection is suspected
• Ultrasound scanning of the pelvis
• Diagnostic laparoscopy, if indicated, to exclude a particular pathology, especially endometriosis

If the findings are normal, reassurance in itself is often sufficient.
MITURITION, FREQUENCY OF

Ben Challacombe

The normal bladder holds around 500 ml. The bladder should store urine until a socially convenient time without urgency or incontinence. Urinary frequency should be less than seven times in 24 hours, although that is, of course, dependent upon fluid intake. In cold weather, most of us notice an increase in urinary frequency. In hot weather, urinary frequency is reduced due to insensible losses, thus reducing urine output. Urinary frequency should be evaluated by asking the patient to fill in a frequency/volume voided chart such that a true indication of normal intake and output can be discovered. Normal daily fluid intake is somewhere between 1500 and 2000 ml, and additional fluid is provided in the food we eat.

Urinary frequency forms part of the symptom complex of the idiopathic detrusor overactivity (IDO, previously called overactive bladder syndrome). The symptom complex includes daytime frequency, nocturia, urgency and urge incontinence. There are many other causes of urinary frequency, some of which are listed in Table M.3.

The majority of pathological conditions within the bladder will be associated with pain or inflammatory symptoms, so before a diagnosis of IDO can be made, a process of appropriate investigation must exclude the more sinister conditions. The frequency chart (Fig. M.14) will demonstrate the relationship between fluid intake and output. Many patients can significantly reduce nocturia by reducing fluid intake after 7 or 8 p.m.

Daytime frequency which is not due to a pathological process can be improved by changes in the volume and timing of fluid intake, and by bladder retraining. Caffeinated drinks cause a physiological diuresis and an increase in urinary frequency. Thus, coffee, tea and sports drinks should be avoided. The normal volume of urine voided lies between 200 ml and 330 ml in young adults, but it tends to reduce with ageing: many older people do not hold more than 150 ml at any time. Thus, before frequency can be attributed to pathology (unless there are other symptoms), these factors must be taken into consideration.

Nocturia is the awakening to pass urine at night. Normal homeostasis ensures that we do not wake at night to pass urine, the effect of antidiuretic hormone on renal homeostasis being in part responsible for the reduction of urine production at night. Normal individuals will wake to pass urine on occasion after drinking larger quantities of fluid prior to going to sleep. However, as we age, more urine is produced at night, and thus there is a likelihood that we will wake at night to void as we reach our 60s and 70s. Other causes of nocturia are as follows:

- Reduced bladder capacity, for the reasons listed above
- Peripheral oedema, particularly dependent oedema in old age or in association with immobility
- Congestive cardiac failure
- Diabetes mellitus and insipidus

These conditions should be treated according to their need.

<table>
<thead>
<tr>
<th>Table M.3 Causes of urinary frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Condition</td>
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<tr>
<td>-------------------------------------</td>
</tr>
<tr>
<td>Urinary tract infection</td>
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<tr>
<td>Urethral syndrome</td>
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<tr>
<td>Radiation cystitis</td>
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<tr>
<td>Prostatic tumours which encroach upon the bladder</td>
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<tr>
<td>Interstitial cystitis</td>
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<tr>
<td>Small functional bladder capacity</td>
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<td>Tumour in the bladder</td>
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<tr>
<td>Bladder stone</td>
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<tr>
<td>Schistosomiasis</td>
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<tr>
<td>Cyclophosphamide cystitis</td>
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<tr>
<td>Chronic retention with overflow</td>
</tr>
<tr>
<td>Bladder instability</td>
</tr>
<tr>
<td>Bladder hyperreflexia due to neurological disease</td>
</tr>
</tbody>
</table>
**Frequency Chart**

Name: ___________________________ Hospital Number: ________________

Start date: _______________________

For 3 consecutive days please record as accurately as possible the number of times you pass urine in the ‘Out’ column and also write the volume of fluid you drink and the times that you drink. Include any episodes of urinary incontinence and the times they occur.

<table>
<thead>
<tr>
<th>Time</th>
<th>Day 1</th>
<th>Day 2</th>
<th>Day 3</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>In</td>
<td>Out</td>
<td>In</td>
</tr>
<tr>
<td>Start</td>
<td>In</td>
<td>Out</td>
<td>In</td>
</tr>
<tr>
<td>6AM</td>
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<tr>
<td>7AM</td>
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<td>Noon</td>
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<td>8PM</td>
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**Figure M.14** Urinary frequency chart.
MONOPLEGIA, UPPER LIMB

**MICTURITION, HESITANCY**

Ben Challacombe

There is normally little delay before the start of micturition in healthy individuals. In potentially socially embarrassing situations, such as in public urinals, there may be a delay in the start of urine flow when trying to void. This delay may be increased substantially in nervous individuals (those with the ‘bashful or shy bladder syndrome’). If there is a significant (non-physiological) delay in the start of urine flow, this is known as urinary hesitancy, and it is one of the classic symptoms of bladder outflow obstruction, which is usually due to benign prostatic enlargement in older men but can also occur in women with bladder hypocontractility. Part of this delay is due to the additional time it takes for the bladder pressure to reach the point at which the outflow resistance is overcome and voiding begins.

Hesitancy will also occur due to detrusor hypo- or acontractility, which may be secondary to long-standing obstruction that has given rise to detrusor dysfunction. This may be the result of anticholinergic medications (oxybutynin, tolterodine or solifenacin) used to treat lower urinary tract symptoms of frequency and urgency. Some over-the-counter medications such as cold medicines and nasal decongestants containing ephedrine or pseudoephedrine may worsen bladder emptying, as may tricyclic antidepressants such as amitriptyline.

Hesitancy may also be the result of neurological disorders that have affected the spinal reflex arc or cauda equina, and it may also follow urinary tract infection or recent surgery. In addition, there is the defined condition, mentioned above – the ‘anxious’ or ‘bashful bladder’, or the ‘executive bladder’ – which is associated with hesitancy and the other urinary symptoms that suggest bladder outflow obstruction. Urinary flow rate analysis followed by formal urodynamic testing should be used whenever there is a doubt over the diagnosis, or where bladder outflow surgery is proposed for a condition in which the diagnosis is not clear.

**MONOPLEGIA, UPPER LIMB**

Mark Kinirons

Monoplegia or monoparesis is defined as loss of motor power involving a single limb. In theory, a monoplegia or monoparesis can result from damage anywhere in the motor pathway between the motor cortex and the muscles of the affected limb. When the degree of weakness is only slight, and associated signs such as increased reflexes are absent, accurate diagnosis may be difficult. For example, localized damage to the hand area of the cortex resulting from a stroke can produce weakness of the hand without any obvious reflex change, mimicking a localized nerve or root lesion. Reaching a diagnosis in such a case requires a careful history and a detailed examination of motor function, with special efforts being made to delineate the extent and pattern of motor weakness.

A patient may complain of weakness in a limb when, in reality, the problem is the result of pain, incoordination, loss of sensation, or akinesia and rigidity, as in Parkinson’s disease. The patient with apraxia of a limb often regards it as weak.

**UPPER MOTOR NEURONE**

A spastic upper limb monoplegia most commonly results from a lesion affecting the motor cortex or the descending corticospinal fibres in the internal capsule. Cerebral infarction, cerebral haemorrhage or tumour is the most likely cause. Muscle atrophy is not a feature, and the appearance of the arm is usually normal. In an acute lesion, muscle tone may be reduced but, given time, spasticity develops and the arm adopts a fixed posture, adducted at the shoulder and flexed at the elbow and wrist. Hand movements are usually more severely affected than shoulder movements. Tendon reflexes are brisk, and contractures may develop. Even when symptoms are apparently confined to the arm, there may be some degree of facial weakness, and the plantar response in the ipsilateral foot may be extensor.

In the patient with a progressive upper motor neurone pattern of weakness, the likeliest cause is an intracerebral tumour. A similar pattern of weakness may occasionally be seen in multiple sclerosis, or rarely in spinal cord tumours. A spastic upper limb monoplegia may be mistaken for the rigid akinetic arm of a patient with Parkinson’s disease. In such cases, the arm moves slowly and is thought by the patient to be weak, yet testing usually reveals no significant loss of power. Similarly, the tendon reflexes in Parkinson’s disease are usually unaffected. Finally, it should be possible to differentiate the rigidity of Parkinson’s disease from the spasticity of an upper motor neurone lesion.

*Functional upper limb monoplegia* is rare. In such cases, the arm appears normal, and muscle tone is not increased. The tendon reflexes remain normal, and the plantar responses are flexor. There is often apparent dense sensory loss in the affected arm, usually with
a sharp line of demarcation at the level of the shoulder. The most difficult cases are those in which there is some genuine weakness with superadded functional overlay.

LOWER MOTOR NEURONE
A flaccid upper limb monoplegia may result from damage to any part of the lower motor neurone between the anterior horn cell and the muscle. Damage to the lower motor neurone is typically associated with muscle weakness, muscle wasting, decrease in the tendon reflexes and fasciculations. Characteristic changes may be found on electromyographic examination.

ANTERIOR HORN CELL DISORDERS
Localized loss of anterior horn cells may produce a flaccid weakness of one or other arm. This may be seen as a common after-effect of poliomyelitis, and the affected limb is often markedly underdeveloped. In motor neurone disease, localized loss of anterior horn cells may produce progressive lower motor neurone weakness in the arm, and this may begin in any of the upper limb muscles. A common pattern is for the weakness to begin in the intrinsic hand muscles, with associated muscle wasting. Prominent fasciculations are common in this disorder. Where there is associated damage to the corticospinal tracts, as in amyotrophic lateral sclerosis, the tendon reflexes are brisk despite obvious muscle wasting and fasciculations. This combination of signs is diagnostic.

Localized lesions in the cervical cord may damage the anterior horn cells and produce a lower motor neurone pattern of weakness. This may be seen in syringomyelia and in spinal cord tumours. These may be diagnosed by the associated signs of sensory loss and corticospinal tract signs in the legs. Ischaemic lesions of the cervical cord may produce discrete loss of anterior horn cells, with localized areas of muscle wasting and weakness. This may be one of the mechanisms whereby the lower motor neurone pattern of weakness develops in cervical spondylosis.

SPINAL MOTOR ROOT DISORDERS
Localized damage to the spinal roots in the cervical region is common as a result of degenerative cervical spine changes (cervical spondylosis). The C5 and C6 segments are most commonly affected, with muscle wasting and weakness being most prominent in the biceps and shoulder girdle muscles. The biceps and brachioradialis reflexes are depressed, and there is often inversion of these reflexes; that is to say, when the biceps tendon is stretched, there is no contraction of the biceps muscle but there is a contraction of the finger flexors, resulting in finger flexion. Extensive cervical spondylosis sometimes produces quite marked wasting and weakness of many of the upper limb muscles. The cause may be recognized by the associated sensory changes and by the presence of pain. Such cases are usually seen in those who have undertaken heavy manual work, and gross changes are usually confined to those over the age of 50 or 60 years. Other causes of damage to spinal motor roots are cervical disc prolapse, trauma to the cervical spine, extramedullary spinal tumours and spinal arachnoiditis.

Herpes zoster is occasionally accompanied by paralysis of one or more muscles within the affected segments. In neuralgic amyotrophy, there is localized muscle weakness and wasting occurring a few weeks after the development of severe pain in the affected limb. There is some debate over the localization of damage to the lower motor neurone in neuralgic amyotrophy; in some cases, the damage may be in the ventral roots, in some in the brachial plexus, and in some in peripheral nerves.

THE BRACHIAL PLEXUS
Brachial plexus damage on one side may usually be recognized by the distribution of muscle weakness and by the associated reflex and sensory changes. Damage may occur as a result of an injury at birth. Downwards traction on the arm in a breech presentation may rupture the upper end of the plexus (C5–C6), with consequent paralysis and atrophy of the spinati, deltoid, brachialis, biceps and brachioradialis. The arm hangs by the side, internally rotated at the shoulder, with the elbow straight and the fingers flexed, the palm of the hand pointed backwards. This is known as Duchenne–Erb paralysis. A similar condition resulting from a fall on the tip of the shoulder is referred to as Erb’s paralysis. In both cases, the motor disturbances are similar to those produced by a lesion of the fifth and sixth segments of the cervical cord, but there are none of the sensory and pyramidal signs that necessarily occur below the level of the lesion in the latter. A second form of obstetric palsy, described as Klumpke’s paralysis, results from injury to the lower cord of the plexus (C8–T1), by traction on the arm in a vertex or shoulder presentation. Atrophic palsy occurs in the intrinsic muscles of the hand and in the flexors of the wrist and fingers. Horner’s syndrome may be present if the roots themselves have been torn. Rarer causes of trauma to the brachial plexus are gunshot wounds or stab wounds. In modern times, perhaps the most common traumatic lesion to the brachial plexus is that occurring as a result of a motorcycle accident.
In such cases, the damage may be not only to the nerves in the brachial plexus, but also to the ventral roots themselves, which may become completely avulsed from the spinal cord.

Upwards spread of an apical bronchial carcinoma may result in erosion into the brachial plexus, most commonly affecting the lower part (Pancoast’s syndrome). Metastatic carcinoma of the breast may produce a similar clinical picture, and this is often difficult to differentiate from post-radiation scarring occurring in a female who has been treated by radiotherapy to the axilla.

THE CERVICAL RIB

The brachial plexus and subclavian vessels cross the uppermost rib as they pass into the axilla. This may be a normal first rib, an ill-formed rudimentary first rib or a cervical rib. The last may be a well-developed ossified structure or a fibrous band (Fig. M.15), but in either case it joins the first thoracic rib lateral to the insertion of the scalenus anterior, at the point where the brachial plexus and vessels pass into the axilla. Such a rib or band tends to angulate the nerves and vessels and to produce friction during respiration and in movements of the shoulder and upper limb, and this in its turn may give rise to pain and paraesthesiae in the ring and little fingers, and atrophic palsy of the small muscles of the hand (see below). However, cervical ribs often give rise to no symptoms at all, and although they are present from birth, symptoms – if they do arise – are uncommon before middle life. They tend to be aggravated by carrying heavy weights and by wearing a heavy overcoat. Friction on the plexus causes atrophy and weakness of the interossei, the thenar and hypothenar muscles, and the long flexors of the fingers and wrist. Fasciculation may be seen in the affected muscles, and slight sensory loss is sometimes found in the medial two fingers and the ulnar border of the hand and forearm. To this essentially neurological picture may be added symptoms and signs of vascular origin: Raynaud’s phenomenon, persistent cyanosis of the hand, diminution or even loss of the radial pulse. The subclavian artery is subjected to friction and to costo-clavicular compression, and this may lead to thrombosis or even to the formation of an aneurysm at the site of compression. If thrombosis occurs, ischaemia gives rise to pallor of the hand, paraesthesiae in all the fingers, claudication of the forearm and hand and, in elderly subjects, necrosis of the fingertips.

Costoclavicular compression can occur even in the absence of a cervical rib. The motor features are similar to those of a cervical rib, but of milder degree, and vascular symptoms tend to predominate. There is aching pain in the shoulder, paraesthesiae in all the fingers, and a feeling of weakness in the limb. These symptoms are intensified by abduction of the arm, by using the hand above the level of the shoulder, as in doing the hair or painting a ceiling, and by recumbency, so that the paraesthesiae are worse at night. These symptoms may come on for the first time in middle-age after a period of prolonged and unwanted physical work, and more commonly in females than males. A troublesome form occurs in the late stages of pregnancy, usually disappearing after parturition and recurring in subsequent pregnancies; it is related to the postural readjustments that take place in pregnancy. Care must be taken not to confuse these symptoms with those due to compression of the median nerve in the carpal tunnel, as in the latter there are also complaints of paraesthesiae in the hand on waking in the morning, with or without wasting of the thenar eminence. In carpal tunnel syndrome, however, the paraesthesiae rarely involve the ulnar part of the hand alone.

PERIPHERAL NERVE DAMAGE

Peripheral nerve damage in the upper limb usually produces localized muscle weakness in the territory supplied by the relevant nerve. Extensive damage most commonly results from lesions in the nerves of the brachial plexus (see above). The term ‘mononeuritis multiplex’ implies multiple peripheral nerve lesions. It may occur in diabetes but is more commonly seen in connective tissue disorders such as polyarteritis nodosa.

Localized weakness in peripheral nerve lesions

The circumflex (axillary) nerve is liable to injury by fractures of the neck of the humerus, by dislocation of the shoulder, by the use of an unpadded crutch, and by penetrating injuries of the axilla and shoulder. The deltoid is paralysed, and there is an area of sensory loss over the proximal half of the lateral aspect of the arm. Paralysis of the radial nerve occurs in fractures of
MONOPLEGIA, UPPER LIMB

the shaft of the humerus, pressure from callus, gunshot wounds of the axilla and arm and, not infrequently, sitting with an arm suspended over the back of a chair. With lesions in the neighbourhood of the shaft of the humerus, the triceps escapes, but there is paralysis of the brachioradialis and of the extensors of the wrist and fingers, with consequent wrist-drop and paralysis of finger extension. There is a somewhat variable loss of cutaneous sensation over the radial border of the forearm and the radial half of the dorsum of the hand, which may be confined to a small patch over the first dorsal interosseous space.

Inability to extend the wrist impairs the mechanical efficiency of the flexors of the fingers, so that the grasp is weakened – a circumstance that may lead to an erroneous diagnosis of a coincident lesion of the median nerve. Wrist-drop is a familiar feature of certain general affections – lead poisoning, leprosy, alcoholic neuritis – but in these conditions the weakness is not wholly confined to the radial distribution.

The posterior interosseous nerve is a branch of the radial nerve in the upper forearm, and it may suffer entrapment or may become the site of a neurofibroma. There is eventually paralysis of the finger extensors, the thumb extensor and abductor of the wrist. The radial extensor is spared so that there is only partial weakness of wrist extension and no wrist-drop. There is no sensory loss. Spontaneous recovery is unlikely, and early exploration is indicated.

Paralysis of the median nerve is usually due to penetrating injuries of the arm or forearm. If the lesion is above the elbow, atrophic palsy involves the pronator teres, flexor carpi radialis, palmaris longus, flexor digitorum sublimis, flexor pollicis longus, pronator quadratus, inner half of flexor digitorum profundus, muscles of the thenar eminence and lateral two lumbricals. Sensory loss and absence of sweating are limited to the median distribution in the hand.

The anterior interosseous nerve is a branch of the median nerve and supplies, essentially, the flexor profundus of the thumb and index finger so that the patient is unable to make a pincer movement. There is no sensory loss. Lesions usually result from entrapment. Spontaneous recovery may occur but, if there has been none after 2–3 months, exploration is indicated.

Compression of the median nerve in the carpal tunnel may cause wasting of the muscles of the thenar eminence, with paralysis of abduction and opposition of the thumb, and cutaneous sensory loss over the thumb, index, middle and radial half of the ring finger. The full-blown picture is rare, but milder degrees of compression occur and can be responsible for acroparaesthesiae in the hand, even when signs are slight or absent.

Carpal tunnel syndrome occurs mostly in women, and usually for no apparent reason. In men, there is likely to be a discoverable cause such as arthritis of the wrist, ganglion at the wrist joint, acromegaly or myxoedema. These causes are not, of course, confined to men, and in women pregnancy is another possible precipitating factor.

The earliest symptom is usually intermittent tingling of the fingers of one hand, often waking the patient from sleep. This tingling may spare the little finger, and is often most prominent in the ring and middle fingers. If the tingling is severe, it is likely to be accompanied by pain in the palm of the hand, sometimes at the elbow or even in the shoulder. As the condition progresses, the patient may find that in the morning the fingers feel swollen and numb. Later, symptoms may occur during the day, and may be brought on by use of the hands. Finally, abnormal signs may appear. These consist of weakness and wasting of the abductor pollicis brevis and sensory impairment within the distribution of the median nerve in the hand, both of which may be quite mild in degree. The condition is relieved by division of the flexor retinaculum at the wrist.

The ulnar nerve is frequently injured by penetrating wounds of the forearm and is particularly liable to compression where it lies behind the medial epicondyle of the humerus. This occurs particularly if the groove in which it lies is shallow, in which case it is subject to recurrent injury, such as in individuals whose occupation entails resting the elbows on a hard table. Cubitus valgus – whether congenital or as a result of a fracture in the region of the elbow – predisposes to this traumatic neuritis. Pain is rare, but paraesthesiae and wasting of the interossei, the hypothenar muscles and the medial two lumbricals cause discomfort and disability. Sensory loss in an ulnar distribution and palpable thickening of the nerve at the elbow afford confirmatory evidence on the nature of the condition. An occupational palsy of the muscles supplied by the deep branch of the ulnar nerve is seen in long-distance cyclists, who lean heavily on the handle-bars, and also in individuals using files, with the instrument held in one hand, and downward pressure exerted by the hypothenar eminence of the other on the end of the file. Weakness and wasting are confined to the interossei, and there is no sensory loss. Ulnar paralysis can occur in leprosy and may be the sole manifestation.
MOUTH, PIGMENTATION OF

Leandros Vassiliou & Mark McGurk

The oral mucous membrane contains melanocytes in the basal layer, as a result of neuroectodermal migration in the fetus, which are similar to those present in the skin. Therefore, any condition that causes abnormal pigmentation in the skin can produce similar changes in the oral cavity, although the effects are usually not as marked. The following are important causes of oral pigmentation.

MELANOTIC NAEVI

These occur less often than in the skin, with the hard palate and the buccal mucosa being the most commonly involved sites. As in the skin, they represent a collection of the normal melanocytes, but instead of being evenly distributed in the basal layer, the cells are aggregated together. Depending on their position in relation to the basement membrane, they give rise to junctional naevi, compound naevi, intramucosal naevi and blue naevi. The lesions are generally small, well circumscribed, macular or slightly raised (Fig. M.16). The majority are pigmented, with varying shades of brown, blue or black. The lesions are twice as common in females as in males and tend to occur in middle-age.

MALIGNANT MELANOMA

This is a rare tumour of the oral mucous membrane with a slight male predominance, again occurring in middle age (Figs M.17 and M.18). It is mostly found in the upper jaw, especially the palate, followed by the gingival mucosa. It is more common in Japanese, Indian and African races, and one-third of cases are preceded by a history of oral pigmentation. As in the skin, any oral pigmented lesion that increases in size, changes its surface characteristics or colour and starts to bleed should be suspected as being a malignant melanoma. Growth of the lesion is followed by destruction of the underlying bone and loosening of the teeth, with rapid spread to the regional lymph nodes. If malignant change is suspected, a wide excision of the lesion should be carried out. Rarely, the mouth may be involved, with secondary deposits from a cutaneous melanoma.

MELANOTIC NEUROECTODERMAL TUMOUR OF INFANCY

This pigmented lesion is invariably noted within the first 6 months of life, the majority occurring in the anterior maxilla. The tumour grows rapidly in size, with underlying bone destruction and displacement of the developing teeth. The correct diagnosis is essential as the tumour is benign and responds well to simple enucleation.

PEUTZ–JEGHER’S SYNDROME

This inherited condition is characterized by intestinal polyposis and melanotic spots of the face and mouth (see Fig. M.19) with the hands and feet also occasionally being affected. Although it is an inherited condition, it is often not suspected until late adulthood.
condition, a family history will not always be found. There are multiple freckles on the face, especially around the mouth (circumoral pigmentation), the eyes and the nose. The polyps in the intestine rarely become malignant, as they are hamartomas in origin.

**ADDISON’S DISEASE**

This condition is caused by bilateral destruction of the suprarenal glands, the most common cause previously being infection with tuberculosis (see Fig. F.6). Today, this condition is usually caused by autoimmune destruction or an opportunist infection in immunodeficient patients, and more recently this has been demonstrated in patients suffering from AIDS. The skin becomes pigmented early on in the disease, especially the exposed areas, while the oral cavity shows patchy melanotic pigmentation, which varies in colour from light brown to black. If this disease is suspected, the diagnosis will be verified by measuring the blood pressure (which is low), the blood urea (which is raised) and the serum sodium (which is lowered). The diagnosis is confirmed by the Synacthen test (the measurement of plasma cortisol in response to an injection of synthetic adrenocorticotrophic hormone (ACTH)).

Pigmentation can also occur with tumours secreting ACTH, the most common being bronchogenic carcinoma, and with Nelson’s syndrome.

**RACIAL**

This is the most common cause of oral pigmentation, which is most prevalent in African and Indian races (Fig. M.20). However, 5 per cent of Caucasian people also show pigmentation of the oral mucosa. The pigment is evenly distributed in the palate, buccal and gingival mucosa. The colour of the pigment is not necessarily related to the colour of the skin.

**AMALGAM TATTOO**

This is a common cause of oral pigmentation, and arises by small amounts of amalgam filling material gaining access to the mucosa via a small abrasion during restorative dental procedures or tooth extraction. This produces small regular or irregular areas of pigmentation in the mucosa, which rarely require treatment (Fig. M.21).

**KAPOSI’S SARCOMA**

The epidemic form of this vascular tumour is a well-recognized complication of HIV infection. Oral lesions are common, typically affecting the palate or gums. The recognition of oral lesions is important, as there is a strong association with concurrent pulmonary or gastrointestinal disease, which frequently requires treatment with chemotherapy.
MOUTH, ULCERS

Leandros Vassiliou & Mark McGurk

The classification of mouth ulcers is outlined in Box M.7.

TRAUMATIC

The diagnosis is usually easy to make because there is a definite history of trauma associated with mastication, ill-fitting dentures or other minor injury to the oral cavity. The ulcers are usually shallow (Fig. M.23) and painful, and heal quickly once the noxious stimulant is removed. Secondary bacterial infection can occasionally occur, causing an abscess or cellulitis.

APHTHOUS ULCERATION

This is the most common form of oral ulceration, and three types are recognized, depending on the size and number of ulcers.

The term minor aphthous ulcer is given to ulcers less than 5 mm in diameter, which occur intermittently as single ulcers, (Fig. M.24) or in crops. The ulcers take

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from 7 to 10 days to heal, and in some patients new ulcers may develop before the original ones have healed, so that they are never without ulceration. This type of ulceration tends to commence in childhood and early life, and the attacks diminish as the patient gets older. They are characteristically found on the buccal mucosa, in the sulcus between the jaws and the cheeks, the ventral aspect of the tongue and the floor of the mouth. This condition is often described as recurrent aphthous stomatitis. The ulcers are round and have an erythematous periphery with a pale central crater.

**Major aphthous ulcers** are larger and more persistent and, in addition to the previous sites, may affect the tongue and the palate (Fig. M.25). They may be up to 1 cm in size and, because of their duration, give concern that the ulcer could be neoplastic.

**Herpetiform ulcers** are the third variant, and here the patient suffers from crops of very numerous small ulcers, which are painful and tend to coalesce into one large irregular area on an erythematous background. Anything from 10 to 100 ulcers may be present at one time.

The cause of aphthous ulceration is unknown, although an autoimmune origin is the most plausible with an inappropriate release of cytotoxic factor ‘heat shock proteins’ to minor stimuli. Some 10 per cent of patients will have an underlying haematological deficiency, especially of vitamin B₁₂, folate or iron; an additional 3 per cent will be suffering from coeliac disease, and a few may have Crohn’s disease. In some female patients, the ulcers are related to the menstrual cycle and will respond to hormone therapy. There is also an association with stress and the cessation of smoking.

Aphthous ulceration is treated by the use of topical steroids and mouthwashes; on occasion, systemic steroids may be required.

**ULCERATION ASSOCIATED WITH OTHER MUCOUS MEMBRANES**

**Behçet’s syndrome**

This syndrome is a systemic illness and consists of recurrent aphthous ulceration with also genital and ocular involvement, in the form of posterior uveitis. The latter may subsequently cause impairment of vision. The disease characteristically affects young men, and there may be associated disease of the skin, joints and nervous system. The cause is unknown, but viral or autoimmune theories have been put forward.

The management of the oral ulceration is the same as for recurrent aphthous ulceration.

**Reiter’s syndrome**

The oral manifestations of this complaint are white circinate lines on an area of erythematous mucosa. These lesions are accompanied by urethritis, arthritis and conjunctivitis. The aetiology is again unknown, but may follow infection with *Mycoplasma* or *Shigella*. Some 10 per cent of patients will show small, non-specific ulcers similar to aphthous ulceration.
MOUTH ULCERATION ASSOCIATED WITH SKIN DISEASE AND CONNECTIVE TISSUE DISORDERS

Lichen planus
Lichen planus is a common condition affecting both the skin and the mouth, although it can affect either in isolation (Fig. M.26). There are several types of oral lichen planus, with the erosive type being characterized by large, irregular areas of mucosal ulceration; the base of the ulcer is often slightly raised, with a covering of white to yellow slough.

Examination of the mouth elsewhere often demonstrates white lacy striations or a desquamative gingivitis. The aetiology of lichen planus is obscure, but it may be mediated by an immunological process with a lymphocytic infiltration, predominantly T-cells, beneath the epithelium. Lichen planus can also be a reaction to liver disease, drugs (e.g. methyldopa, gold salts or antimalarials), or in the graft-versus-host reaction (bone marrow transplantation).

Pemphigus and pemphigoid
Pemphigus and pemphigoid both may produce oral lesions, which commence as bullae when the epithelium separates from the basal layer in pemphigus, and when the epithelium and basal layers separate from the underlying mesoderm in pemphigoid (Fig. M.27). As a result, the bullae in pemphigus are far more fragile and rupture quickly, whereas the bullae in pemphigoid are more resilient.

In pemphigus, there are circulating autoantibodies against the intercellular attachments of the squamous epithelium, and the level is an indication of the severity of the disease. Immunofluorescence shows deposits of IgG and C3 binding to desmosomes, which is the area of contact between epithelial cells. In pemphigoid, circulating autoantibodies are not detectable, but immuno-fluorescence studies show IgG and C3 at the basal membrane or dermo-epidermal junction.

Pemphigus is a serious vesiculobullous mucocutaneous condition that mostly affects woman in the 40-to 50-year age group. If untreated, the condition may be fatal. The mouth may often be affected first, with small bullae or widespread ulceration and loss of the oral epithelium. As the bullae are intraepithelial, they rupture easily, leaving what appears to be a thin layer of tissue paper over an underlying erosion. The diagnosis is made by biopsy (preferably of an intact bulla) and immunofluorescence studies. Treatment is by the use of systemic steroids, with replacement of fluid and protein in the acute phase.

*Mucous membrane pemphigoid* is a disease of the elderly, seen in females more than males. It affects the oral mucous membrane, and predominantly the conjunctiva of the eyes are involved. Anogenital lesions may occur, and minor involvement of the skin may also be noted. Because the bullae are more rigid, they tend not to enlarge and rupture late. Once ruptured, they leave areas of irregular ulceration, which are accompanied by considerable scarring. The oesophagus and nasopharynx may also be involved, but the most significant aspect of the disease is conjunctival fibrosis leading to visual disturbance.

Examination of the mouth will demonstrate several intact or ruptured bullae during the active phase of the disease, and the gingivae may be severely affected with a desquamative gingivitis. Treatment is with topical steroids, although occasionally systemic steroids may be necessary.

*Bullous erythema multiforme*
Bullous erythema multiforme, or Stevens–Johnson syndrome, is the more severe form of erythema multiforme, invariably involving the oral cavity, and...
this can be the predominant feature of the attack. The appearance is dramatic because of the severe oral ulceration and the blood-stained and crusted lips (Fig. M.28). It tends to affect children and young adults, and is probably immunologically mediated, via exposure to microorganisms (e.g. herpes simplex and Mycoplasma), or from drugs, typically the sulphonamides, non-steroidal anti-inflammatory agents, phenytoin and penicillin.

Examination of the skin will demonstrate either extensive erythema or a macular rash with target lesions exhibiting central bullae formation or ulceration. There is conjunctivitis, leading to corneal ulceration. In the mouth, diffuse inflammation leads to vesicle formation followed by widespread erosions and haemorrhage. Epistaxis commonly occurs. Treatment of the minor case is with topical steroids, but the more severe attack may require systemic therapy. Tetracycline antibiotics should be given if infection with Mycoplasma is suspected.

Angina bullosa haemorrhagica
This condition of unknown origin is characterized by the spontaneous formation of blood-filled blisters within the mouth, which can develop very rapidly. They may occur while eating, and generally involve the soft palate. The individual is often alarmed by this condition, and may develop a sensation of choking. These blisters normally rupture within 24 hours, leaving an ulcer that heals spontaneously. Although similarities exist between this condition and mucous membrane pemphigoid, immunofluorescence tests are negative. Blood coagulation studies are normal.

Lupus
Approximately 20 per cent of patients with discoid and systemic lupus erythematosus will show oral ulceration. There may be areas of erythema with erosions and, in some areas, these may resemble lichen planus due to minor striae formation. Frank ulceration may also be present.

The differential diagnosis may be difficult to make, but here the lesions often occur on the hard palate, which is rare in lichen planus. The diagnosis is made by biopsy and immunofluorescence studies. Antinuclear antibodies should be sought in the serum, and a history of arthritis and skin rashes should be elicited, particularly an erythematous rash of the face in the butterfly distribution.

BLOOD DYSCRASIAS
Blood dyscrasias are discussed elsewhere (see GUMS, BLEEDING, p. 228), but ulceration of the gingivae is also common, caused by acute local bacterial infection secondary to the abnormal white cell function. It should be remembered that the ulceration may also be due to acute bacterial ulcerative gingivitis or acute viral herpetic gingivostomatitis that has arisen because of a blood dyscrasia as the primary cause.

ULCERATION ASSOCIATED WITH GASTROINTESTINAL DISEASES
Although coeliac disease, ulcerative colitis and Crohn's disease can be associated with recurrent aphthous ulceration, these conditions may also show distinctive oral signs (Fig. M.29). Crohn's disease characteristically produces a 'cobblestone' thickening of the buccal mucosa, with hyperplastic folds and fissuring. Painful ulcers, which are slow to heal, may also be present. Inflammatory bowel disease may also produce the condition of pyostomatitis vegetans, which
is characterized by soft hyperplastic mucosal folds between which fissures and ulcers may form. The ulceration associated with coeliac disease and gluten hypersensitivity is known as dermatitis herpetiformis. The skin is affected by an itchy rash, while in the mouth erythematous areas may appear or extensive erosions. Linear IgA disease may also cause oral ulceration.

INFECTION

Bacterial infections

Acute ulcerative gingivitis

Bacterial infection of the oral mucous membrane is rare in normal circumstances, the most important example being acute ulcerative gingivitis, which is associated with large numbers of Vincent’s organisms (Treponema vincentii and Fusiformis fusiformis). There is haemorrhage, inflammation and the formation of painful, shallow ulcers at the crest of the gingival margin. This eventually leads to destruction and flattening of the interdental papillae. Infection is associated with pre-existing periodontal disease and also any condition reducing host immunity. Patients with leukaemia or AIDS, and those receiving chemotherapy, are therefore all susceptible, and infection may follow an episode of acute herpetic stomatitis.

Acute ulcerative gingivitis in the severely debilitated patient may progress to the destructive condition of cancrum oris (see GUMS, BLEEDING, p. 227). This is rare in the developed world, apart from those patients who are immunosuppressed, but still occurs in developing countries as a result of malnutrition and severe viral infections (e.g. herpes and measles). Small areas of gangrene appear in the lips, cheeks or other oral structures, which rapidly progress to larger areas of slough and extensive loss of facial tissue.

Syphilis

The oral cavity may be involved rarely, during the primary stage, to produce a chancre on the lips or the tip of the tongue. Secondary lesions known as the ‘mucous patch’ are now seldom seen because of modern effective treatment. Here, there is an erosion in the mucous membrane of a few centimetres with a yellowish slough surrounded by erythema. When these areas coalesce, they give an irregularly shaped ulcer known as a ‘snail-track’ ulcer. There has been resurgence in the incidence of this condition and the diagnosis should always be considered in the appropriate circumstances. It was always known as the great mimic and can present in unusual ways.

The tertiary stage of syphilis, now rarely seen, produces the gumma, which is a deeply punched-out ulcer caused by central necrosis. These may typically affect the tongue or the palate and, in the latter, perforation will produce a central oronasal fistula.

Tuberculosis

With the successful treatment of tuberculosis, oral lesions are rare but, when they do occur, they are usually found in the tongue and lips. The ulcer typically shows undermined edges and a granulating floor. The mode of infection is thought to be expectoration of tubercle bacilli from a primary focus in the lungs, which then become implanted in the oral cavity.

Fungal infections (Candida)

This opportunistic infection (thrush) is caused by the yeast-like fungus Candida albicans, which invades the epithelium, causing erythema of the epithelium and a yellow soft plaque that can be easily removed (Fig. M.30). This leaves an area of haemorrhagic mucosa or ulceration.

A high proportion of patients suffering with HIV infection will commonly develop an infection with Candida. This is known as erythematous and pseudomembranous candidosis, which are descriptive terms to describe the appearance of this infection.

Viral infections

The majority of acute infections of the oral mucosa are viral in origin. They tend to affect the younger age groups, and they are normally associated with constitutional symptoms of fever, malaise and enlarged and tender cervical lymph nodes.

Herpes simplex

The primary infection invariably involves the mouth, with many small vesicles, approximately 2 mm in diameter, of regular shape and size (Fig. M.31).
The mouth is generally inflamed and, in some areas, the ulcers coalesce to produce irregular raw erosions and a yellow slough.

The gingival mucosa is red and swollen and may bleed, even in the absence of ulceration. Herpetic stomatitis may be an opportunist infection, and can be severe in immunocompromised patients and patients with AIDS. The lesion of reactivation is known as ‘secondary herpes’. This reactivation can be brought on by exposure to sunlight, menstruation, trauma and stress, and particularly immunosuppression. The virus, which has been quiescent in the trigeminal ganglion, becomes active once more, to produce a ‘cold sore’ or herpes labialis (Fig. M.32) at the mucocutaneous junction of the lip or the nostril. The ulceration occurs in the distribution of the infected nerve. Treatment is by the use of aciclovir.

Herpes zoster
Herpes zoster causes chickenpox in the patient who has not been exposed to the virus, and mouth ulcers are common in this condition. The virus again remains quiescent in the trigeminal ganglion, and on reactivation causes painful ulceration within the exact anatomical distribution of the nerve (Fig. M.33). The ulceration may be preceded by pain or a disturbance in sensation in the same area.

This condition is known as herpes zoster or shingles and, on occasions, it can be an indication of a more serious underlying disease. The picture once more is of erythema, small regular ulcers and pain within the distribution of the nerve affected. If the ophthalmic branch is involved, care needs to be given to the cornea. The infection may be complicated by postherpetic neuralgia. Treatment again is with the antiviral agent aciclovir.

Epstein–Barr virus
Infectious mononucleosis is caused by the Epstein–Barr virus (EBV). There may be characteristic petechiae of the soft palate and pharynx, which are often considered diagnostic.

The patient has a sore throat, the fauces and palate become inflamed and oedematous, and the lymph nodes are enlarged. Occasionally, there is extensive ulceration of the fauces. The diagnosis is achieved by the identification of atypical mononuclear cells in the peripheral blood, the Paul–Bunnell test and by EBV serology. Ampicillin should not be given for the sore throat as this exacerbates the condition.

Coxsackie virus
Mouth ulcers may arise from infection with the Coxsackie A virus, which causes a febrile illness, lymphadenopathy, and ulcers and intense erythema of the soft palate.
Epidemics – particularly among children with the Coxsackie A virus – give rise to hand-foot-and-mouth disease. This is a highly infectious condition, characterized by small vesicles, leading to ulcers on the hands, on the feet and in the mouth. These infections generally are trivial conditions and are self-limiting.

CANCER
Squamous cell carcinoma is the most common malignant tumour of the oral cavity and is associated with excessive smoking and alcohol consumption. In the Indian subcontinent, it is associated with the chewing of a betel nut quid with added tobacco and other chemicals (Fig. M.34).

The clinical picture of oral carcinoma is very varied, but there are two important patterns. In the mouth, the cancer always produces an ulcer unless there has been previous surgery or radiotherapy. But, in the oropharynx, the cancer can burrow beneath the mucolymphoid tissue and be hidden from sight. This can lead to delayed diagnosis and medico-legal complaints. Usually, there is a friable mass arising in the mucous membrane with a rough, irregular surface, which bleeds easily. There is also usually deep, irregular central ulceration with an infected slough at its base. Radiographs may show associated bone erosion. The early lesion may be more difficult to diagnose, showing only an area of erythema or hyperkeratosis, slight roughening of the mucosa and shallow ulceration.

Unfortunately, many oral carcinomas are still diagnosed late. Any ulcer that does not heal within 3 weeks, or does not obviously fall into one of the other categories, should be submitted for a biopsy.

MUSCULAR ATROPHY
Mark Kinirons
Muscular atrophy occurs for a number of reasons. Patients present complaining of various symptoms such as wasting muscles or, most commonly, weakness of specific muscles leading to an inability to perform certain tasks.

DAMAGE TO THE LOWER MOTOR NEURONE
Lesions of the lower motor neurone are responsible for the majority of cases of muscular atrophy encountered in clinical practice. The muscles are weak, flaccid and wasted. Fasciculation is seen. Electromyography shows fibrillation potentials on mechanical stimulation by the exploring needle, and spontaneous fibrillation and fasciculation potentials at rest. The tendon reflexes are depressed or abolished in the affected part unless, as in motor neurone disease, there is a coincident pyramidal lesion, in which case they usually remain brisk as long as there is any power of contraction left in the muscles concerned. It is convenient to classify the causes of this form of atrophy according to whether the lesion is in the anterior horn cells, the motor roots or the peripheral nerves.

DISUSE
Disuse of a limb gives rise to extreme degrees of wasting. The best example is provided by the results of prolonged immobilization for fractures of the long bones. It also occurs in muscles acting upon a painful or ankylosed joint, and in rheumatoid arthritis. Voluntary disuse, as practised by some Eastern cults, can lead to extreme ‘withering’ of the entire limb – for example, when penance demands that the arm be held above the head in perpetuity.

VASCULAR DISEASE
Arterial occlusion is variable in its effects. In tourniquet paralysis, surgical ligation of a major vessel, thrombosis or embolism of the great vessels, the degree of wasting depends on the efficiency of the collateral circulation. In many instances, the muscles become fibrosed and hard rather than wasted and flabby. In other cases, there is a true atrophy of muscles due to ischaemia of the motor nerves, as in some cases of Buerger’s disease and in polyarteritis nodosa. A special variety of ischaemic palsy, known as Volkmann’s contracture, is the result of fractures in
the region of the elbow. Oedema and haemorrhage into the soft tissues interfere with the circulation distal to the elbow and cause fibrosis of the flexors of the wrist and fingers, and of the intrinsic muscles of the hand. The affected muscles are hard in consistency, and wasting is comparatively slight owing to replacement with fibrous tissue.

GENERAL BODILY WASTING
Muscular atrophy is often seen in general wasting of the tissues due to chronic or subacute disease such as tuberculosis, malignant disease, neglected or undiagnosed diabetes mellitus, hyperthyroidism, malaria, hookworm infestation, conditions associated with chronic diarrhoea and anorexia nervosa. It is also seen in old age.

ANTERIOR HORN CELLS
Acute poliomyelitis used to be the most common cause of anterior horn cell damage, but the incidence of this has been dramatically reduced. The muscular atrophy seen in late cases of polio is often very striking and is localized to particular groups of muscles. Localized damage to anterior horn cells may occur at the site of trauma from fracture-dislocation of the spine, or as a result of backward protrusion of an intervertebral disc in the cervical or thoracic regions. Spinal tumours may produce a similar effect. In these instances, atrophy is confined to the muscles that are innervated by the affected portion of the cord. Other intraspinal lesions that may produce muscular atrophy are syringomyelia, haematomyelia and intramedullary tumours such as ependymomas and astrocytomas; such cases usually have associated sensory loss and other signs of spinal cord damage. Motor neurone disease is a common cause of muscle atrophy in adults. It is prominent in the progressive muscular atrophy form of the disease, and is also seen in the so-called ‘amyotrophic lateral sclerosis’ form. Fasciculations are common in affected muscles in this condition. In the bulbar form, wasting of the tongue may be obvious. The rare syphilitic atrophy may mimic motor neurone disease, but the tendon reflexes are depressed and serological tests will confirm the diagnosis. Infarction of a localized area of spinal cord consequent upon thromboembolism of a branch of the anterior spinal artery may cause localized damage to anterior horn cells, with segmental muscle wasting. This may occur in atheromatous disease, in meningeovascular syphilis or in association with arteriovenous malformations or tumours of the spinal cord.

MOTOR ROOTS
Damage to a single motor root seldom causes much atrophy since most muscles are supplied by several roots. An exception is the first thoracic root, which supplies the intrinsic muscles of the hand (Fig. M.35). Damage to multiple spinal roots may be seen in spinal lesions below the level of the first lumbar vertebra, producing damage to the cauda equina. Apart from muscle weakness and wasting, there will often be associated sensory loss and sphincteric disturbance; pain is often a prominent symptom. Common causes of cauda equina damage include lumbar spondylosis, lumbar canal stenosis, primary and secondary spinal tumours, and prolapsed lumbar intervertebral discs. More localized damage to spinal roots may be seen in spondylosis, and muscle atrophy in the upper limbs may be a feature of cervical spondylosis.

THE PERIPHERAL NERVES
Proximal damage to the peripheral nerves may occur at their origin in the brachial or lumbosacral plexus. Malignant infiltration of the brachial plexus typically affects the lower part of the plexus, thereby affecting the nerve supply to the intrinsic hand muscles. Malignant infiltration of the lumbosacral plexus may occur but is less common. Pain is a prominent feature. Traumatic avulsion of the plexus in the upper limb is all too common as a consequence of motorcycle injuries, and in some instances the avulsion occurs not at the level of the plexus but at the level of the motor roots. Traumatic damage to the lumbosacral plexus is less common. Localized injuries to the plexus or peripheral nerves may occur as a result of stab wounds, gunshot wounds, and fractures and dislocations in the vicinity of motor nerves.

Compression on motor nerves may produce local damage (so-called ‘entrapment neuropathies’). Common sites of damage are the radial nerve in...
the upper arm, the lower cord of the brachial plexus between the first rib and a cervical rib causing thoracic outlet syndrome, the ulnar nerve at the level of the elbow, the median nerve at the level of the carpal tunnel, and the lateral popliteal nerve at the level of the neck of the fibula. The causes of peripheral neuropathies are listed in Box M.8.

Rates of progression vary considerably, but the common features are those of diffuse involvement of multiple peripheral nerves. In general, the distal parts of the limbs are affected first, with muscle weakness and wasting being most obvious in the lower part of the legs and hands (Fig. M.36). Signs are usually symmetrical, although some types of peripheral neuropathy tend to involve motor fibres more than sensory fibres. However, there is usually an associated distal sensory loss, which is typically in ‘stocking and glove’ distribution. Tendon reflexes are almost invariably depressed or absent.

**MUSCLE DISORDERS**

Muscular atrophy, although not usually a prominent feature in muscle disease, may be seen in any form of myopathy. (For a classification of myopathies, see Box M.9.) Congenital absence of a muscle may occur, and the most common deficiency is the sternocostal portion of pectoralis major. Rupture of a muscle or its tendon may lead to localized atrophy.

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**Box M.8 Causes of peripheral neuropathy**

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<thead>
<tr>
<th>Inherited neuropathies</th>
<th>Uraemic</th>
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<tbody>
<tr>
<td>• Mixed sensorimotor neuropathies</td>
<td>• Beri-beri</td>
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<tr>
<td>– Idiopathic</td>
<td>• Diabetic</td>
</tr>
<tr>
<td>– Metabolic</td>
<td>• Hypothyroid</td>
</tr>
<tr>
<td>• Sensory neuropathies</td>
<td>• Connective tissue</td>
</tr>
</tbody>
</table>

**Acute acquired neuropathies**

| Idiopathic |
| Porphyria |
| Toxic |
| Diphtheritic |

**Subacute acquired neuropathies**

| Deficiency states |
| Heavy metals |
| Drug intoxication |
| Uraemic |
| Diabetic |
| Arteritic |

**Chronic acquired neuropathies**

| Carcinoma |
| Paraproteinaemia |

**Relapsing neuropathies**

| Idiopathic |
| Porphyria |

**Mononeuropathies**

| Pressure |
| Trauma |
| Idiopathic |
| Serum/post-vaccinal |
| Herpes zoster |
| Neoplastic |
| Leprosy |
| Radiation |
| Diphtheritic |
| Mononeuritis multiplex |
| – Arteritis |
| – Diabetes |

**Box M.9 A classification of myopathies**

| Inflammatory disease of muscle |
| – Polymyositis |
| – Dermatomyositis |

**Infective myopathies**

| Trichinosis |
| Toxoplasmosis |
| Virus infections (e.g. Coxsackie B virus) |

**Muscular dystrophies**

| Duchenne |
| Becker |
| Facioscapulohumeral |
| Limb girdle |
| Oculopharyngeal dystrophy |
| Myotonic dystrophy |

**Metabolic myopathies**

| Glycogen storage myopathies |
| Carnitine deficiency (and other defects of fatty acid metabolism) |

**Endocrine myopathies**

| Adrenal dysfunction |
| Thyroid dysfunction |

**Spinal muscular atrophies**

| Although such conditions are not strictly disorders of muscle, they are included because of the frequent difficulty in differentiating them from the true myopathies |
MUSCULAR HYPERTROPHY

Mark Kinirons

There is considerable variation in the size of individual skeletal muscles, and the assessment of hypertrophy or enlargement may in some instances be very difficult. For example, the normal calf muscles in a leg that has gross wasting of the quadriceps may appear hypertrophied simply because of the relative disproportion.

Exercising muscles can cause them to hypertrophy; this occurs as a result of increase in muscle fibre size secondary to an increase in the number of myofibrils per fibre. Such physiological hypertrophy may be generalized, as in body-builders, or localized, as in those who repeatedly use a particular limb, for example tennis players.

The syndrome of hemi-hypertrophy is associated with diffuse enlargement of all the tissues of one half of the body, occasionally affecting only the face or one limb. In this condition, the aetiology of which is unknown, the muscles, subcutaneous tissues and bones all appear to enlarge.

Pathological muscle hypertrophy may be classified as in Box M.10.

MULTIPLE MUSCLE ENLARGEMENT

Enlargement of muscles is commonly seen in Duchenne and Becker dystrophies, and occasionally in limb-girdle dystrophy. The most commonly enlarged muscle is the gastrocnemius, although hypertrophy of the infraspinatus, deltoid, triceps, quadriceps or gluteus muscles may be encountered. The term ‘pseudo-hypertrophy’ has been used in reference to Duchenne dystrophy, since, in the later stages of the disease, the muscles become weak but remain enlarged because of replacement of the fibres by fat and connective tissues. Often the pseudo-hypertrophic muscles will have a rather characteristic firm or doughy feeling.

Rapid enlargement of the muscles has been described in spinal muscular atrophy, polymyositis and cysticercosis, and in certain families with malignant hyperpyrexia.

Diffuse hypertrophy of the muscles may occur in myotonia congenita and probably represents work hypertrophy associated with continuous muscle contraction. This produces the so-called ‘infant Hercules’ appearance. In hypothyroidism, hypertrophy of the muscles is more common in children than in adults. The calves, thighs, hands, neck, tongue and face may all enlarge and feel firm or indurated. Often the patient will complain of pain and stiffness, and this may be accompanied by proximal muscle weakness. In Cornelia de Lange’s syndrome, hypertrophy of the muscles is associated with congenital athetosis and mental retardation.

LOCALIZED MUSCLE ENLARGEMENT

This is most commonly seen in the biceps muscle as a result of rupture of the muscle fibres or tearing of the long head. It may also be seen as a result of a muscle haemorrhage, muscle tumour such as rhabdomyosarcoma, angiomma, desmoid or metastatic lesion, or as a result of an infective process such as granulomatous disease or pyogenic abscess. Localized enlargement may occur as a result of trauma leading to myositis ossificans.

MUSCULAR PAIN

Mark Kinirons

Pain in muscles is one of the most common of all complaints, and it may occur either as an ache or as a stiffness or cramp. These two types of muscle pain are usually distinguishable, and their causes are different.

ACHING PAINS

Aches and pains in muscles may be transient, recurrent or persistent. They may be classified as follows.

Normal reactions

Fatigue, anxiety and particularly depression may be associated with muscle aches and pains. Few healthy people go through an average day, whatever their occupation or activity, without occasional feelings of muscular discomfort. These are usually suppressed or ignored, but in abnormal emotional states, they may assume unnatural proportions, and become significant and sometimes distressing symptoms. Muscle aches become more common as individuals become increasingly aware of ageing. These aches may be aggravated by postural strain, often resulting from the adoption of fixed positions over prolonged periods.
Injuries
After unduly heavy exertion, muscles may become painful for hours or even days, presumably as a result of small local injuries, tears or haemorrhages, in muscles themselves or at muscle insertions. Injured fibres, particularly at points of attachment to bone, may be acutely painful and tender, as in tennis or golfer’s elbow, and ligaments may be completely torn across even in the absence of extreme exertion. The largest tendon in the body, the Achilles tendon, is particularly liable to rupture in those engaged in sporting activities such as squash or cricket. Too rapid a leaping out of bed has been known to rupture both tendons. The supraspinatus muscle and tendon and the quadriceps are perhaps the most frequently injured; a rupture of the long head of biceps, the rhomboïd, major or minor, pectoralis major, rectus abdominis, trapezius, levator scapuli, latissimus dorsi or almost any other muscle or tendon may occur, and may mislead the unwary diagnostician. It is important that muscle pain in the chest or abdomen should not be mistaken for pain arising from visceral sources. An unusual cause of muscle pain that may be overlooked is spontaneous haemorrhage of the type that may occur in individuals with haemophilia or in those on anticoagulant therapy.

Referred pain
Pains apparently arising in muscles are often referred from inflammatory or degenerative disease in nearby joints. Pains in the thigh muscles, for instance, may result from hip joint disease, and muscle pains in the shoulder girdle, chest or abdomen from degenerative changes in the cervical, dorsal or lumbar spine. Disease or injury of the ligaments may be referred to muscles; injections of hypertonic saline into the interspinous ligaments, for instance, may cause pains in muscles several centimetres away. The cause of the extremely common fibromyalgia in muscles in the scapular and shoulder regions is still inadequately explained, but the pain is undoubtedly often referred from adjacent spinal joints and ligaments, allegedly triggered in many cases by cold or damp. These ‘fibrositic’ pains may sometimes be the first signs of an arthropathy such as rheumatoid arthritis, or rarely a disseminated connective tissue disorder such as systemic lupus erythematosus or polyarteritis nodosa, or, more commonly in elderly subjects, polymyalgia rheumatica (see pp. 168, 331 and 429).

Pains of vascular origin
The classic example of ischaemic muscle pain is the intermittent claudication experienced on exertion by victims of occlusive arterial disease in the lower limbs. Inflammatory arterial diseases (e.g. polymyalgia rheumatica, giant-cell arteritis and polyarteritis nodosa) may also produce tenderness in muscle, but not pain. Whenever pain is a presenting symptom, consideration of the blood supply is vital in assessment.

Muscle disorders
Several inflammatory disorders of muscle are commonly associated with muscle pain and tenderness, and these should be considered in any patient who complains of myalgia. In polymyositis and dermatomyositis, over 50 per cent of sufferers will describe pain as well as muscle weakness. The pain is usually a deep aching within the muscles, and it is often aggravated by activity. The muscles may be swollen as well as tender. Similar symptoms are seen in a variety of other connective tissue diseases when there is an associated inflammatory myopathy, for example rheumatoid arthritis, Sjögren’s syndrome, systemic lupus erythematosus, polyarteritis nodosa, scleroderma and mixed connective tissue disease.

Several epidemics of influenza due to type A or B virus have been described in which there was a rather acute onset of severe myalgia lasting 1 or 2 weeks, with creatine kinase often elevated many times above normal. In Bornholm’s disease, the acute onset of chest and abdominal pain, chiefly at the costal margins and in the subcostal region, is usually the first symptom. Affected muscles are tender to pressure, and pain is induced by muscle contraction. The acute pain and fever last several days, and after initial recovery, one or more relapses are not uncommon. Care must be taken not to confuse one of these relatively benign and reversible syndromes with polymyositis of acute onset.

An unusual type of muscle disease may result from eating undercooked pork infected with the larvae of Trichinella spiralis. Muscle pain and stiffness occur particularly in the masseter. There may be associated malaise, fever, periorbital oedema, skin rash and petechial haemorrhages.

Polymyalgia rheumatica should always be considered in older patients in their 60s and 70s who complain of aching muscles and stiffness in the shoulder and hip girdle muscles. The ache tends to be worse at night, and morning stiffness is a prominent feature.

Neuropathic disorders
Pains in the muscles may be marked in the early stages of polyomylitis and are common in Guillain–Barré syndrome. In acute brachial neuritis or neuralgic amyotrophy, severe aching pain in one or both shoulders or arms may precede the development of localized muscle weakness. In diabetic proximal
neuropathy, pain often accompanies atrophy of the quadriceps or sometimes other muscles. Other neuropathies associated with severe pain include those seen with alcoholism, arsenic poisoning, polyarteritis and porphyria.

Endocrine causes
The muscles may ache in hypothyroidism, and also in acromegaly and hyperparathyroidism. Most endocrine disorders, however, are more commonly associated with weakness than with pain.

Drugs
Severe muscle pains may follow the intramuscular injection of suxamethonium, a muscle relaxant used in anaesthesia; these pains can occur 1–5 days after anaesthesia, and are worse if there is an earlier resumption of physical activity. A number of drugs may cause an acute or subacute necrotizing myopathy, which may be associated with myoglobinuria, sometimes leading to acute renal failure (Box M.11).

Psychogenic
Aching pain in muscles is a common symptom in depression and anxiety, and it may also be seen in hysteria or in patients who are frankly malingering.

MUSCLE STIFFNESS OR CRAMPS
Ordinary muscle cramps or painful involuntary contractions of muscle are experienced in almost everyone at some time in their lives, and in most instances the diagnosis presents no difficulty. Benign muscle cramps most commonly occur after unaccustomed exercise and are usually self-limiting. In some patients, however, these cramps can be persistent and quite incapacitating.

Pathological cramps can be produced by an abnormality anywhere along the final common pathway, including the anterior horn cell, peripheral nerve, neuromuscular junction and muscle membrane. Exertional cramps may be seen in a variety of disorders, including cauda equina ischaemia from lumbar canal stenosis, and inborn errors of muscle metabolism such as McArdle’s disease or the lipid storage myopathies. In myophosphorylase deficiency and phosphofructokinase deficiency, a true contracture occurs with exercise, which may be accompanied by myoglobinuria.

Several of the cramping disorders are characterized by myotonia, where the patient observes a delayed relaxation after even a single voluntary contraction. Muscle cramps and spasms may occur either spontaneously or after exercise in a variety of biochemical disturbances. They are most characteristically seen in tetany, which may occur with hypocalcaemia, alkalosis or hypomagnesaemia.

Several disorders are characterized by an almost continuous state of muscle activity; these include tetanus, strychnine poisoning and the bite of the American black widow spider. A number of rare neuromuscular disorders are associated with persistent contraction of the muscles at rest. These include the symptoms of myokymia, continuous muscle fibre activity and the so-called ‘stiff man’ syndrome.

A separate and important group of muscle cramp syndromes are the so-called ‘professional’ cramps or repetitive strain injury; the most common of these is ‘writer’s cramp’. However, similar localized muscle cramps have been described in typists, telephone operators, painters, tailors, seamstresses, pianists, flautists, violinists, cellists, harpists, drummers, piano players, blacksmiths, file-makers and watchmakers. There is considerable debate over whether these are occupational neuroses or localized dystonias.

MUSCULAR TONE
Mark Kinirons
The tone of a muscle can be regarded as its resistance to passive stretching. This can only be assessed when the muscle is moved; it cannot be judged on the basis of palpation. Postural tone is that state of partial contraction of certain muscles that is needed to maintain the posture of the body.

At the moment a muscle is stretched, receptors in the muscle concerned – particularly in the muscle spindles – transmit afferent stimuli, and reflex partial contraction of the muscle results. The responses to momentary stretching are responsible for the tendon jerks; the responses to more prolonged stretching elicit more complex responses, such as tonic contraction.

Forceful continued contraction of a group of muscles – for example, clenching one hand – temporarily causes an increased flow of afferent impulses in the sensory fibres from the spindles. This is associated with

<table>
<thead>
<tr>
<th>Box M.11 Drugs capable of causing painful myopathy</th>
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<tr>
<td>• Suxamethonium</td>
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<tr>
<td>• Clofibrate</td>
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<td>• Epsilon-aminocaproic acid</td>
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<td>• Amphetamine</td>
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</table>
exaggeration of the tendon reflexes resulting from increased alpha-neuronal discharge. This is known as ‘reinforcement’, which can be helpful in eliciting depressed reflexes.

Muscle tone is normally regulated by reticulospinal fibres, which accompany the pyramidal tract through the spinal cord, and which have an inhibitory effect upon the stretch reflex. This inhibition balances the facilitatory impulses conveyed by the pontine reticulospinal and lateral vestibulospinal pathways. These in turn are influenced by multisynaptic reflex arcs traversing the cerebellum, basal ganglia and brainstem.

ALTERATIONS IN MUSCLE TONE

Rigidity
In this form of increased muscle tone, the muscles are continuously or intermittently tense. The increased resistance to passive movement has an even or uniform quality throughout the range of movement of the limb, like that noted when bending a lead pipe. Rigidity is present in all muscle groups, both flexor and extensor, but it tends to be more prominent in those which maintain a flexed posture.

A particular type of rigidity, often encountered in Parkinson’s disease, has distinctive characteristics and is described as ‘cogwheel’. When the hypertonic muscle is stretched, a ratchet-like resistance is felt.

Rigidity is a prominent feature of many extrapyramidal diseases such as Parkinson’s disease, Wilson’s disease, dystonia musculorum deformans, multiple system atrophy and basal ganglia calcification.

A special type of variable resistance to passive movement is that in which the patient seems unable to relax. This is sometimes called ‘gegenhalten’, and it is seen in diseases of the frontal lobes. A similar difficulty may be observed in children.

The pathophysiology of rigidity is not fully understood, although it is generally agreed that it results from lesions of the nigrostriatal system. It can often be relieved by dopamine, dopamine agonists or stereotactic lesions in the ventrolateral thalamus.

Spasticity
This is a form of increased tone resulting from lesions of the pyramidal pathways and the closely associated reticulospinal pathways. The stretch reflexes are released from descending inhibitory influences, and there is increased excitability of the fusimotor neurones and alpha motor neurones. The resistance to passive stretch of muscles affected by spasticity often has a distinctive quality. It may be particularly severe initially and then tend to give way – hence the descriptive term ‘clasp-knife rigidity’. The hyperactivity of the tendon reflexes is often accompanied by clonus in which sustained stretch of a muscle evokes repetitive contraction and relaxation.

Decerebrate rigidity
A lesion at the midbrain, at or about the level of the superior colliculus, releases the brainstem, cerebellum and spinal cord from cerebral control. This results in strong continuous contraction in extensor groups of muscles. In such cases, all four limbs are rigidly extended, the back is arched, and there may be neck retraction.

Although the experimental production of decerebrate rigidity by Sherrington appeared to have a specific anatomical substrate, this does not hold quite true in man; for example, unilateral decerebrate rigidity is not uncommonly seen in the early stages of acute cerebral infarction in the territory of the middle cerebral artery.

HYPOTONIA

Hypotonia, or flaccidity, implies a reduction in tone. It may be seen in cerebral or spinal shock resulting from acute damage to the brain or spinal cord. It is a common manifestation of cerebellar disease, and in this situation it is thought to result from diminished gamma efferent activity. It also occurs whenever a lesion interrupts the afferent or efferent pathway of the spinal reflex arc. It is thus seen in damage to the anterior horn cells, spinal roots and peripheral nerves, but it is not a particular feature of muscle disease.

Minor degrees of loss of tone may be difficult to differentiate from normal inherent muscle tone. Gross flaccidity is usually obvious. In the upper limbs, lesser degrees of hypotonia can be elicited by asking the patient to hold the arms out horizontally. Tapping the forearms when hypotonia is present will be associated with slow recoil and a wide arm swing.

MYOCLONUS

David Werring

Myoclonus describes the shock-like contractions of a group of muscles, irregular in rhythm and amplitude, and usually asynchronous and asymmetrical. It can be a useful clue to the localization and type of neurological disease. Myoclonus can be classified according to the part of the nervous system thought to generate it: namely cortical; cortico-subcortical; subcortical (non-segmental); segmental; and
peripheral. If myoclonus affects the whole body at once, it can be termed generalized. Another widely used way to classify myoclonus is according to clinical features and causes (Box M.12); this includes physiological, essential, epileptic and symptomatic (secondary) forms. The classification of myoclonus is important as each subtype has different diagnostic implications. Myoclonus is often stimulus- or activity-sensitive, in which case it can be termed reflex or action myoclonus, respectively.

**PHYSIOLOGICAL MYOCLONUS**

This form of myoclonus occurs in normal individuals. A variety of sleep jerks occur in sleep or transition between sleep phases (e.g. when drifting off to sleep). Partial myoclonic jerks are usually multifocal, in distal muscles. Generalized myoclonic jerks affect trunk and proximal muscles. Periodic movements of sleep are repetitive dorsiflexion of the toes and feet, sometimes with flexion of the knees and hips.

**ESSENTIAL MYOCLONUS**

In essential myoclonus there are either no or only minor additional features. Hereditary essential myoclonus typically begins before age 20, is autosomal dominant with variable penetrance and is benign. Myoclonus usually occurs throughout the arms and axial muscles. It is worse with muscle activity and much improved by drinking alcohol. Dystonia is commonly a feature, and then the syndrome is called myoclonus-dystonia. Genetic causes have been found, including mutations in the epsilon-sarcoglycan or DYT1 genes. Sporadic essential myoclonus is a heterogeneous group, and some cases may actually be cases of missed family history.

**PALATAL MYOCLONUS**

This rhythmic form of clonus is sometimes classified as a tremor because it is continuous. There is an essential form, in which there is rhythmic contraction of the tensor veli palatini muscles (up to about 7 Hz), causing a repetitive auditory click. There is no known pathological cause. The second and more common (symptomatic) form involves the levator palatini muscles (up to about 3 Hz), with rapid elevation of the uvula and palate. There may be oscillopsia and cerebellar signs, and it persists in sleep. The condition is seen when lesions interrupt the tegmental tracts that connect the midbrain nuclei to the inferior olivary complex (Mollaret’s triangle). The cause may be vascular, demyelinating, neoplastic or traumatic.

**EPILEPTIC MYOCLONUS**

This is myoclonus in patients with various types of epilepsy, with prominent seizures. Fragments of epilepsy, with prominent seizures. Fragments of

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**Box M.12 Classification of myoclonus**


**Physiological (occurring in normal subjects)**
- Sleep jerks
- Anxiety-induced
- Exercise-induced
- Infantile benign myoclonus associated with feeding
- Hiccoughs

**Essential**
- Hereditary (autosomal dominant)
- Sporadic
- Palatal with no brainstem lesion

**Myoclonic epilepsies (associated with seizures)**
- Fragments of epilepsy: isolated epileptic myoclonic jerks; epilepsy partialis continua; stimulus-sensitive; photosensitive; absence with minor myoclonus or myoclonic absences
- Childhood myoclonic epilepsies: infantile spasms (West syndrome); severe infantile myoclonic epilepsy (Dravet syndrome); benign myoclonic epilepsy of infancy; Lennox–Gastaut syndrome; myoclonic-astatic epilepsy; juvenile myoclonic epilepsy
- Familial cortical myoclonic tremor with epilepsy

**Symptomatic (secondary) not defined by seizures**
- Progressive myoclonic epilepsy (Unverricht–Lundborg disease, also known as Baltic myoclonus)
- Storage diseases: Lafora body disease; GM2 gangliosidosis; Tay–Sachs disease; Gaucher disease; Krabbe leukodystrophy; neuronal ceroid lipofuscinosis; sialidosis type 1 and 2
- Spinocerebellar degenerations: progressive myoclonic ataxia (Ramsay Hunt cerebellar syndrome; Friedreich’s ataxia; ataxia telangiectasia; other
- Basal ganglia degenerations: Wilson disease; torsion dystonia; pantothenate kinase-associated neurodegeneration; progressive supranuclear palsy; Huntington’s disease; Parkinson’s disease; multiple system atrophy; corticobasal degeneration; dentatorubral pallidolysian atrophy (DRPLA)
- Dementias: Alzheimer’s disease; Creutzfeldt–Jakob disease; dementia with Lewy bodies; frontotemporal lobar degeneration; Rett syndrome
- Diffuse infectious/postinfectious/inflammatory disorders: subacute sclerosing panencephalitis; encephalitis lethargica; herpes simplex encephalitis; other
- Metabolic: hyperthyroidism; hepatic failure, renal failure, electrolyte disturbances
- Toxic/drug-induced: bisulfide or heavy metal toxicity; l-dopa; psychotropic drugs including tricyclics; anticonvulsants; other
- Physical: post-hypoxic (Lance–Adams); post-trauma, electric shock; decompression syndrome
- Focal CNS injury: stroke, tumour, trauma, multiple sclerosis; other
- Malabsorption: Whipple’s disease; coeliac disease
- Opsoclonus–myoclonus syndrome: paraneoplastic; infectious; idiopathic; other

(Continued)
brief myoclonic jerks may also be a feature of typical absence seizures in adolescence.

**SYMPTOMATIC (SECONDARY) MYOCLONUS**

Myoclonus can be a symptom of a myriad of neurological and non-neurological disorders (see Box M.12).

**Progressive myoclonic epilepsies**

Progressive myoclonic epilepsies are characterized by severe and progressive myoclonus, generalized tonic–clonic seizures, dementia and ataxia. There are many causes but the commonest ones are Lafora body disease, myoclonic epilepsy with ragged red fibres (MERRF), neuronal ceroid lipofuscinosis, dentatorubral pallidoluysian atrophy, and storage diseases including Gaucher disease. *Lafora body disease* is familial (autosomal recessive), has an onset in late childhood or adolescence, and is characterized by polyglucosan-Schiff-positive inclusion bodies in the brain, liver, muscle or skin (eccrine sweat glands). *Myoclonic epilepsy with ragged red fibres (MERRF)* is maternally inherited (transmitted by mitochondrial DNA) and may be diagnosed by increased serum and cerebrospinal fluid lactate and ‘ragged-red’ fibres on muscle biopsy. Neuronal ceroid lipofuscinosis (*Batten disease*) presents with seizures, myoclonus, dementia and visual loss (in the childhood forms), and it is characterized by curvilinear inclusion bodies (lipofuscin) in the brain, eccrine glands, muscles and gut. In the adult-onset patient with progressive myoclonic epilepsy, *dentatorubral pallidoluysian atrophy (DRPLA)* should be considered, especially (but not exclusively) in Japanese patients. This disorder has a wide phenotype, which may include dystonia or a bradykinetic–rigid syndrome. *Unverricht–Lundborg disease* is also familial and is characterized by stimulus-sensitive myoclonus, tonic–clonic seizures, EEG abnormalities (paroxysmal generalized spike-and-wave activity and photosensitivity), ataxia and mild dementia, with an onset at 5–15 years of age. Disorders of lysosomal storage can also cause progressive myoclonic epilepsy. In these conditions, there is a deficiency of one or other enzymes responsible for the breakdown of intracellular lipids (sphingolipids) or other complex molecules such as mucolipids or glycoproteins. These molecules thus accumulate in neurones, causing progressive and widespread neurological dysfunction. *Sialidosis* is a storage disorder associated with a ‘cherry-red spot’ on fundoscopy and dysmorphic facial features. *Tay–Sachs disease* is a disorder of sphingolipid storage with an infantile onset, marked startle response, profound
motor and cognitive delay, spasticity, visual loss, cherry-red spot, seizures and death age 3–5 years. The late-childhood-onset form of Gaucher disease is associated with supranuclear gaze palsies, ataxia and splenomegaly; characteristic abnormal histiocytes (Gaucher cells) are seen on marrow, liver or spleen biopsy.

**Spinocerebellar degenerations**

Progressive myoclonic ataxia (Ramsay Hunt cerebellar syndrome). This syndrome is distinguished from progressive myoclonic epilepsy by the relative lack of dementia and prominent ataxia, and it has a number of causes. These include mitochondrial encephalomyopathy, coeliac disease, late-onset neuronal ceroid lipofuscinosis, biotin-responsive encephalopathy, adult Gaucher disease, action myoclonus–renal failure syndrome, neurodegenerative diseases including spinocerebellar degenerations, olivopontocerebellar atrophy and DRPLA. Hypoxic damage to the brain causes polymyoclonus in the acute phase, but as recovery occurs, a striking form of intention or action myoclonus becomes apparent, which is always associated with cerebellar ataxia. This is termed post-anoxic or Lance–Adams myoclonus. When an attempt is made to move a limb, large amplitude myoclonic jerks occur that are distinct from classical intention tremor. This disabling syndrome may improve with clonazepam or valproate treatment. Opsoclonus–myoclonus consists of dramatic, chaotic eye movements due to ocular clonus (or marked ocular dysmetria), associated with polymyoclonus and ataxia. It is a non-metastatic, immunologically-mediated manifestation of cancer (i.e. a paraneoplastic syndrome). In children, it is usually associated with medulloblastoma, but in adults it is mainly associated with breast and small-cell lung cancer. Rarer associations are with melanoma and lymphoma. The relevant antineuronal antibodies can be detected in specialized laboratories to confirm the diagnosis.

**Myoclonus with dementia**

In middle and later life, polymyoclonus with dementia must raise the suspicion of Creutzfeldt–Jakob disease or Alzheimer's disease. Much rarer causes include Whipple's disease of the central nervous system (which causes the rare but pathognomonic movement disorder of oculomasticatory myoarhythmia), corticobasal degeneration (in which an asymmetrical bradykinetic–rigid syndrome with alien limb are typically seen), DRPLA and AIDS dementia. In children with prolonged polymyoclonus and encephalopathy, subacute sclerosing panencephalitis, a rare entity related to measles virus infection, should be considered. Selected causes of metabolic and toxic myoclonus, which may present at any age, are listed in Box M.12.

**SPINAL MYOCLONUS**

Myoclonus may arise from pathology in the spinal cord rather than the cortex, cerebellum or brainstem. Important causes are herpes zoster myelopathy, spinal trauma and the myelopathy associated with spinal cord demyelination in multiple sclerosis.
NAILS, AFFECTIONS OF
Barry Monk

The nails are absent or hypoplastic from birth in cases of ectodermal dysplasia; this may be a familial trait, and a search should be made for associated abnormalities of the hair, the teeth and – most importantly – the sweat glands; in such individuals, heat or exercise may lead to collapse or sudden death. In another genetic disorder, epidermolysis bullosa, repeated blistering of the fingertips may lead to scarring and loss of the nails. Other genetic syndromes involving the nails include Darier’s disease and pachyonychia congenita. Spoon-shaped nails, koilonychia, may also be congenital, but in acquired koilonychias there is likely to be a background of chronic iron deficiency. The nails may be shed spontaneously, usually after some major febrile illness; more commonly, there is a temporary disruption to nail growth, producing a transverse depression running across each nail (Beau’s lines) (Fig. N.1) that gradually grows out. Other manifestations of systemic disease including clubbing of the fingers (see FINGERS, CLUBBED, p. 200), whitening of the nails (leuconychia) in chronic hepatic disease, and red half-moons, which are said to be a feature of some cases of cardiac failure. By contrast, white dots and patches within the nailplate, leuconychia striata and leuconychia punctata, are of no pathological significance. In the yellow nail syndrome (Fig. N.2) the nails take on a yellowed or greeny-yellow colour, grow excessively slowly, and are excessively curved; lymphoedema is often found in association, as are various pulmonary abnormalities including bronchiectasis, and benign pleural effusions. Splinter haemorrhages under the nailplate used to be regarded as a marker of infective endocarditis, but they are a non-specific sign, often provoked by trauma. Nailfold infarcts may be a feature of many vasculitic disorders.

Nail changes are a feature of many inflammatory skin disorders. In psoriasis, the nails are affected in up to 50 per cent of cases (Fig. N.3); changes include pitting of the surface of the nail, onycholysis (separation of the distal nailplate, producing the appearance of whitening of the distal nail), discoloration of the nail, or a general thickening and friability of the nail. None of these features is totally specific for psoriasis. Thus, onycholysis may be seen in cases of thyrotoxicosis, or alone as an idiopathic phenomenon. Pitting may be an occasional feature of alopecia areata. In dermatitis of the fingers, there is quite commonly a rather non-specific ridging of the nailplate, while in lichen planus, inflammation around the nailfold may eventuate in scarring destruction of the nail (pterygium).

Thickening and crumbling of the nails may be a feature of dermatophyte fungus infection, and this is
more common on the toenails than the fingernails. *Trichophyton rubrum* and *Trichophyton mentagrophytes* are the most common causative species (Fig. N.4). The changes may be indistinguishable from psoriasis, so it is always wise to confirm the diagnosis by fungal culture before embarking on treatment. However, in their earliest stages, fungal infection may merely present with a localized whitening of the nailplate (*superficial white mycosis*).

In chronic paronychia, which is due to combined candidal and bacterial infection, pockets form under the posterior nailfolds, which will admit the tip of a probe up to 3 mm. Very occasionally, pus may be expressed from these pockets, from which cultures of the yeast may be made. In profile, the raised nailfolds over the pockets have a characteristic bolster-like appearance. The changes in the nailplate are secondary. This disease occurs in those who are obliged to keep their hands wet for long periods, notably bar staff, fishmongers and those with ‘wet’ occupations. It is much more common in women than in men.

Longitudinal ridging more often has no apparent cause. Brittleness (*onychorrhexis*) or softness of the nails with splitting or terracing of the free edge is more common in women than in men. The cause is unknown, but repeated wetting and drying may be a factor of importance. It has been thought to be aggravated by the use of nail varnish and varnish removers, but many cases occur in women who do not use nail polish. Hypertrophy of the nails (*onychauxis*) occurs more often on the toes and may lead to onychogryphosis (Fig. N.5).

*Onychia,* or inflammation of the nail, is usually septic in origin, but it may be the result of trauma or contact with irritants at work. It usually terminates in shedding of the affected nail. Acute *paronychia,* or whitlow, is usually obvious (Fig. N.6), but a primary chancre of the finger may sometimes imitate it. *Trauma* is responsible for many nail deformities. Single injuries produce haematomas that may lead to temporary nail loss or to permanent splits if the nail matrix is injured (Fig. N.7).
Occasionally, an unrecognized injury that leads to a haematoma may lead to the mistaken impression of a subungual malignant melanoma. Periungual warts may distort nail growth, as may an adjacent myxoid cyst. A solitary periungual wart must be distinguished from a squamous cell carcinoma or Bowen's disease.

**NAPKIN (PERINEAL) ERUPTIONS**

Dipak Kanabar

Napkin dermatitis is very common in young infants, particularly those who wear occlusive plastic pants and nappies (diapers). It is usually responsive to emollients and home remedies. Most examples are caused by irritation of the skin due to prolonged contact with urine or faeces and are favoured by infrequent nappy changes. The convexities are affected by shiny red plaques, sometimes eroded, which spare the flexural creases. If the flexural creases are the main site of inflammation, secondary candidiasis should be suspected, and a search made at the edge of the rash for typical 'satellite pustules'. Infantile seborrhoeic dermatitis is an acute self-limiting red scaly eruption in which the napkin area is usually severely affected. The condition begins with cradle cap and spreads to the face, chest, back and limb flexures, as well as the napkin area. Despite the widespread eruption, infants show little sign of discomfort. This is in marked contrast to those infants with atopic dermatitis, who may also have a napkin eruption, but who are distressed by itch and sleep poorly.

Irritable children with erosions and papules around their napkin areas should be carefully examined for scabies; other common sites for burrows are the anterior axillary folds, wrists, finger webs, ankles, palms and soles. Nursing mothers may have burrows around their nipples as well as the usual sites. Molluscum contagiosum causes persistent irritating papules around the anogenital area in both children and adults. Patients are frequently atopic, and individual lesions may heal spontaneously, sometimes with a considerable purulent reaction or a stark halo of eczema.

Earlier editions of this book considered nappy rash seen as a manifestation of congenital syphilis. Affected infants are irritable, feed poorly and have a purulent often bloody nasal discharge. Perianal and perioral erosions and rhagades are prominent and highly infectious. At 2 months of age, the napkin area as well as the palms and soles may be affected by a new pinkish rash resembling he rash of secondary syphilis in adults.

In the extremely rare acrodermatitis enteropathica, zinc absorption is deficient. Affected infants have diarrhoea and characteristic peri-orificial erosions, which may spread onto the limbs, sometimes in a livedoid pattern. Similar lesions can occur in zinc-deficient premature babies. If a napkin eruption becomes purpuric, Letterer-Siwe disease (histiocytosis) should be suspected. Other flexures, the gums and the external auditory meatuses should be inspected for lesions.

White patches in the perineum can be due to vitiligo, which may begin in childhood and has a predilection for genital skin.

Lichen sclerosus of the vulva (white spot disease) is rare in infancy, but young children are affected. The physical signs of a porcelain-white, scarred, purpuric vulva may lead to suspicions of molestation.

Hyperpigmentation is most commonly due to the Mongolian spot, a reliable marker of Negroid or Asiatic parentage, which causes considerable and extensive bluish pigmentation of the sacral napkin area and is due to dermal dendritic pigment cells.

In adults, the perineum may be involved in numerous dermatoses with scaly eruptions, e.g. tinea (see Fig. S.12), erythrasma, candidiasis or psoriasis. The area can be infested in pediculosis pubis, as well as scabies. Molluscum contagiosum can cause irritating papular lesions. Venereal warts are increasing in frequency, and these can be a marker of other sexually transmitted infections or cervical intra-epithelial neoplasia in women. Syphilitic chancre occurs more commonly within the anal canal or vagina, but rarely syphilitic condyloma acuminata may be seen on the perineum. In pruritus ani, lichenification from scratching and rubbing will be seen. There may be a complicating contact dermatitis due to one of the topical remedies employed (e.g. local anaesthetics, local antibiotics or lanolin).
NASAL DEFORMITY
Michael Gleeson

Nasal deformity can be congenital or developmental but is most often acquired. Abnormalities of either the bony or cartilaginous nasal skeleton contribute to the deformity.

Minor degrees of septal deformity are extremely common and are thought to be the result of undetected minor injuries during birth or early childhood. These may compromise nasal airflow but rarely cause significant external nasal deformity. However, in some individuals, deformity may develop during adolescence. One theory is that this is caused by unequal growth of the nasal skeleton as a result of previous trauma.

The most common cause of an acquired deformity is a simple, displaced, fracture of the nasal bones. More severe blunt injuries to the nose are often associated with fractures and severe distortion and deviation of the nasal septum. In this situation, there is a visible displacement of the nasal tip. From a practical standpoint, nasal fracture should be diagnosed clinically rather than radiologically, as non-displaced fractures are irrelevant, and plain X-rays can be extremely unreliable and difficult to interpret.

*Saddle nose deformities* are usually acquired as a result of destruction of the septal cartilage (Fig. N.8).

\[\text{Figure N.8 Saddle nose. The cause of this was a submucous resection operation for a septal haematoma.}\]

Septal abscesses or haematoma, arteritides (e.g. Wegener’s disease or relapsing polychondritis) or, nowadays rarely, congenital syphilis may cause this appearance. It is also a recognized complication of excessive resection of septal cartilage during surgery and cocaine abuse.

It must be recognized that perception of appearance is very subjective and can even be influenced by fashion. Humps, bulbous, broad, raised or drooping nasal tips are inherent in some. These features may detract from the patient’s potential appearance and can have a profound effect on personality. Rhinoplasty can go some way to correcting or changing these deformities in selected cases and significantly improve the patient’s quality of life (Fig. N.9). Unfortunately, there are a few patients for whom surgery can never provide a satisfactory solution, and they require psychiatric help.

NASAL DISCHARGE
Michael Gleeson

The causes of nasal discharge are listed in Box N.1.

CONGENITAL CAUSES

*Choanal atresia*, by blocking the posterior choana, prevents the normal nasal mucus stream from reaching the pharynx. A mucoid anterior nasal discharge results, and intermittent infections ensure that the discharge is mucopurulent at times.

INFECTIVE CAUSES

*Acute viral infection*, the common cold and the prodromal stages of many infectious fevers give rise to a clear mucoid discharge. The discharge becomes increasingly purulent as secondary bacterial infection follows. Persistent purulent discharge, either unilateral or bilateral, is seen in sinusitis. Careful inspection...
of the nasal passages may reveal the origin of the discharge and indicate the sinus or sinuses involved. The maxillary sinuses open into the middle meatus, as do the anterior ethmoid sinuses. The posterior ethmoid and sphenoid sinuses drain more posteriorly. About 10 per cent of maxillary sinus infections have a dental cause, for example periapical infection. If pus aspirated from the maxillary sinus is foetid, a dental cause should be suspected. Sometimes, the pus becomes inspissated, giving rise to the so-called ‘caseous sinusitis’ or ‘rhinitis’. This is particularly common with fungal infection. The material is whitish, cheesy and foul smelling. Even worse are the crusts and dry discharge of an atrophic rhinitis. In these patients, the nasal fossae are unduly wide, and the mucosa thin and devoid of mucus cells. The especially disgusting odour is called ‘ozaena’.

Pathogenic fungi and yeasts are a common cause of nasal infection and discharge in tropical countries but nowadays these infections are occasionally seen in this country in immigrants and travellers returning from exotic places.

To summarize the nasal symptoms of the main fungal diseases:

• **Rhinosporidiosis** predominantly affects the nasal mucosa, where the characteristic lesion is a bleeding polyp containing the sporangium, from which spores spread via the lymphatics (Fig. N.10). This condition chiefly affects the peoples of Sri Lanka, Bangladesh and India.

• **Phymomycoses** cause serious disease, often starting with granulomatous lesions in the nose and considerable mucoid discharge. This fungus is also found in the tropics.

• **Aspergillus infection** is sometimes contracted from captive birds. It is characterized by a watery, mouldy smelling discharge, and a greyish membrane on the mucosa.

• **Actinomycosis** rarely affects the sinuses and nose. There is a woody mass and multiple sinuses from which the pus exudes.

• **Candida albicans** is found commonly in the mouth and occasionally in the nose of the young and those in a poor state of general health. It forms white patches that can be removed without bleeding. Many patients start with nasal discharge and bleeding. The diagnosis is made by identifying the fungus in scrapings, or by biopsy.

• **Secondary syphilis** affects the nose, causing a simple catarrhal rhinitis. Other accompanying lesions usually suggest the diagnosis. Tertiary syphilitic gummas were, at one time, relatively common and frequently affected the nose, with destruction of bone and cartilage. Secondary infection causes offensive discharge, and there is bleeding, often with the later development of an atrophic rhinitis.

• **Lupus vulgaris** is the most common tuberculous infection of the nose. There is nasal discharge, and the typical lesion – a reddish, firm nodule – is found at the anterior end of the nasal septum. The septum may perforate.

• **Leprosy** affects the nose as almost the earliest sign of the disease. Nodular thickening of the mucosa with inflammation and obstruction are associated with discharge. Later, perforation of the septum and destruction of tissue allows for secondary infection and very offensive discharge.

• **Rhinoscleroma** is a progressive granuloma beginning in the nose in an atrophic form.

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<tr>
<th>Box N.1 Conditions causing nasal discharge</th>
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<tbody>
<tr>
<td><strong>Congenital</strong></td>
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<tr>
<td>• Age-related (senile) cataract</td>
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<tr>
<td>• Cataract associated with ocular disease</td>
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<td>• Choanal atresia</td>
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<tr>
<td><strong>Infective</strong></td>
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<td>• Acute rhinitis</td>
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<td>• Rhinosinusitis</td>
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<td>• Caseous rhinitis</td>
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<td>• Atrophic rhinitis</td>
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<td>• Chronic infective granulomas</td>
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<td>– Syphilis</td>
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<td>– Leprosy</td>
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<tr>
<td><strong>Miscellaneous</strong></td>
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<tr>
<td>• Wegener’s granulomatosis</td>
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<td>• Old age</td>
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**Figure N.10** The typical appearance of rhinosporidiosis, seen here in a young girl from Bangladesh.
with ozaena. Nodules form later, and there is considerable scarring. The diagnosis is made by biopsy.

- AIDS is associated with a watery nasal discharge that is frequently found concomitantly in this disease.

**FOREIGN BODIES**

A unilateral nasal discharge in an infant or child is pathognomonic of a foreign body, often a piece of foam rubber or paper inserted into the nasal fossa. Small hard objects may remain in the nose for some time before symptoms develop. Long-standing foreign bodies calcify to form rhinoliths. These are associated with chronic discharge and halitosis.

**TRAUMA**

A clear watery discharge following a head injury suggests a fracture of the anterior cranial fossa or cribiform plate and a cerebrospinal fluid (CSF) leak. Such a discharge will increase on leaning the head forward, or on compression of the jugular veins. The presence of sugar in this fluid confirms that it is CSF. This can be further confirmed by serological tests for beta-transferrin.

**ATOPIЯ**

Copious clear mucoid nasal discharge, almost as freely flowing as water, is present in allergic rhinitis and may be associated with violent attacks of sneezing, lacrimation and conjunctival injection. The diagnosis is confirmed by the history and by radioallergosorbent tests (RASTs) and paper radioimmunosorbent tests (PRISTs). Nasal allergy is liable to be confused with less non-specific vasomotor rhinitis, which is also very common. Numerous provocative factors are present in this troublesome condition, including changes of environmental temperature and humidity, mechanical irritation from dusts and vapours, psychological factors, pregnancy and drug reactions. Patients with vasomotor instability not infrequently carry boxes of paper tissues. The diagnosis is made after excluding nasal allergy.

**NEOPLASTIC CAUSES**

*Malignant nasal disease* (squamous cell carcinoma, adenocarcinoma or olfactory neuroblastoma) is usually heralded by progressive unilateral obstruction, followed shortly by nasal discharge. This is often clear to start with, but it later becomes blood-stained, offensive and thick. The growth may be in the nasal fossa, one or other of the sinuses or the nasopharynx. This sequence of events should prompt the clinician to obtain a computed tomography scan of the patient, and to examine the nose under anaesthetic so that a biopsy can be obtained.

The so-called ‘midline granuloma’ is in fact a T-cell lymphoma that presents as a slowly progressing ulceration of the face, starting in the region of the nose and pre-maxilla. The localized destruction of tissue produces a blood-stained nasal discharge.

**MISCELLANEOUS CAUSES**

*Wegener’s granulomatosis* commonly involves the nose and usually presents with symptoms referable to the paranasal sinuses. It also affects the kidneys, lungs and lower respiratory tract. Patients present with malaise, severe facial pain, nasal obstruction and discharge spotted with blood.

*Senile rhinorrhoea* is thought to be a form of vasomotor rhinitis. It is a common and sometimes distressing condition for the elderly to bear and is not easy to treat. There are no physical signs apart from a watery nasal drip.

**NASAL OBSTRUCTION**

Michael Gleeson

Nasal obstruction – the inability to breathe through one or both nostrils – is a very common complaint, and the reason for referral to an ENT outpatient clinic. The causes of nasal obstruction and discharge are listed in Box N.2.

*Choanal atresia* may be membranous or bony, unilateral or bilateral. Nasal respiration is innate and

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**Box N.2 Causes of nasal obstruction and discharge**

<table>
<thead>
<tr>
<th>Congenital</th>
<th>Acquired</th>
</tr>
</thead>
<tbody>
<tr>
<td>Choanal atresia</td>
<td>Physical obstruction</td>
</tr>
<tr>
<td>• Bilateral</td>
<td>• Adenoids</td>
</tr>
<tr>
<td>• Unilateral</td>
<td>• Deviated nasal septum</td>
</tr>
<tr>
<td>Acquired</td>
<td>• Concha bullosa</td>
</tr>
<tr>
<td>Physical obstruction</td>
<td>Trauma</td>
</tr>
<tr>
<td>• Adenoids</td>
<td>• Haematoma and abscess of septum</td>
</tr>
<tr>
<td>• Deviated nasal septum</td>
<td>• Inflammation</td>
</tr>
<tr>
<td>• Concha bullosa</td>
<td>• Viral, bacterial or fungal rhinitis</td>
</tr>
<tr>
<td>Trauma</td>
<td>• Allergic rhinitis</td>
</tr>
<tr>
<td>• Haematoma and abscess of septum</td>
<td>• Nasal polyps</td>
</tr>
<tr>
<td>• Inflammation</td>
<td>• Rhinitis medicamentosa</td>
</tr>
<tr>
<td>• Viral, bacterial or fungal rhinitis</td>
<td>Autonomic imbalance</td>
</tr>
<tr>
<td>• Allergic rhinitis</td>
<td>• Vasomotor rhinitis</td>
</tr>
<tr>
<td>• Nasal polyps</td>
<td>Neoplastic</td>
</tr>
<tr>
<td>• Rhinitis medicamentosa</td>
<td>Epithelial</td>
</tr>
<tr>
<td>• Autonomic imbalance</td>
<td>• Papilloma</td>
</tr>
<tr>
<td>• Vasomotor rhinitis</td>
<td>• Inverted papilloma (Ringerz tumour)</td>
</tr>
<tr>
<td>• Schneiderian polyp</td>
<td>• Schneiderian polyp</td>
</tr>
<tr>
<td>• Squamous cell carcinoma</td>
<td>• Squamous cell carcinoma</td>
</tr>
<tr>
<td>• Transitional cell carcinoma</td>
<td>• Transitional cell carcinoma</td>
</tr>
<tr>
<td>• Non-epidermoid</td>
<td>• Adenocarcinoma</td>
</tr>
<tr>
<td>• Adenocarcinoma</td>
<td>• Adenocystic carcinoma</td>
</tr>
<tr>
<td>• Neuro-ectodermal</td>
<td>Neuro-ectodermal</td>
</tr>
<tr>
<td>• Olfactory neuroblastoma</td>
<td>Olfactory neuroblastoma</td>
</tr>
<tr>
<td>• Vascular</td>
<td>Vascular</td>
</tr>
<tr>
<td>• Angiofibroma</td>
<td>• Angiofibroma</td>
</tr>
<tr>
<td>• Connective tissue tumours</td>
<td>• Connective tissue tumours</td>
</tr>
<tr>
<td>• Fibroma</td>
<td>• Fibroma</td>
</tr>
<tr>
<td>• Fibrous dysplasia</td>
<td>• Fibrous dysplasia</td>
</tr>
<tr>
<td>• Fibrosarcoma</td>
<td>• Fibrosarcoma</td>
</tr>
<tr>
<td>• Chondrosarcoma</td>
<td>• Chondrosarcoma</td>
</tr>
<tr>
<td>• Miscellaneous</td>
<td>Miscellaneous</td>
</tr>
<tr>
<td>• Lymphoma</td>
<td>• Lymphoma</td>
</tr>
<tr>
<td>• Plasmacytoma</td>
<td>• Plasmacytoma</td>
</tr>
</tbody>
</table>
therefore, in bilateral cases, the obstruction must be bypassed immediately after birth by placement of an oropharyngeal airway. Surgical correction of the abnormality is undertaken in the first few days of life. Unilateral choanal atresia is sometimes not discovered until later. Babies breathe continuously while feeding; hence, those with unilateral choanal atresia are unable to do so when the patent nostril is blocked by their mother’s breast. Consequently, they suckle better from one breast than the other.

Deviations of the nasal septum are extremely common, and present in up to 30 per cent of normal individuals. It has been speculated that these deviations are caused by birth trauma, but there is little evidence to support this contention. The majority of deviations are acquired by trauma after birth as a result of sports injuries, falls or physical conflicts. Some of these deviations are severe enough to cause either unilateral or bilateral obstruction. In some, the sinus outflow tracts are impaired, and recurrent sinus infection ensues. For these patients, septoplasty is a simple and effective remedy. Septoplasty alone is unlikely to restore a normal nasal contour or profile in those who have acquired their deviations traumatically. For these, a septrhinoplasty (see NASAL DEFORMITY, p. 442) will be more appropriate. Septal deviation can be seen with a nasal speculum. It will often be found that the inferior turbinate in the concavity of the deviation will have hypertrophied, thus increasing the obstruction. It is important to remember that the spur of the deflected septum may obscure other pathology behind.

Adenoid enlargement is probably the most common cause of nasal obstruction in childhood. These lymphoid masses usually regress in the first decade of life. In a few children, hypertrophy can cause near-complete obstruction and result in loud snoring and obstructive sleep apnoea. Adenoidectomy is curative.

In recent years, it has been realized that pneumatization of the anterior end of the middle turbinate can be a cause of nasal obstruction and recurrent sinus infections. This condition, termed ‘concha bullosa’, is easily eradicated by a small endoscopic procedure (Fig. N.11).

A subperichondrial haematoma of the nasal septum may develop spontaneously, but it is more commonly caused by blunt trauma or inadequate packing after nasal surgery. It may be either unilateral or bilateral, and presents as a tender soft-tissue swelling within the nose. The haematoma should be evacuated by removal of a patch of mucosa, as the pressure may cause cartilage necrosis and later a saddle nose deformity. Infection sometimes intervenes to produce a septal abscess; this is an extremely painful condition and is almost always followed by significant cosmetic deformity (Fig. N.12).

Swelling of the mucosa, sufficient to block the airway, may be caused by acute or chronic infection (rhinitis). The common cold is the most simple and frequent example of this. Chronic rhinitis is most often associated with sinusitis, which needs to be addressed in order to relieve the obstructive symptoms.

In certain parts of the world, specific bacterial infections (rhinosporidiosis and rhinoscleroma) are a relatively common cause of protracted nasal obstruction. These conditions require long-term antimicrobial therapy and, in the case of rhinosporidiosis, surgical resection of the obstructive growth.

Allergic rhinitis is the most common manifestation of type I allergy, affecting up to 20 per cent of the population in Western countries. Both acute and chronic forms are recognized. In the acute form – hayfever – the

![Figure N.11](a) Coronal computed tomography (CT) scan of the sinuses showing aeration of both middle turbinates, known as concha bullosa (arrows). (b) CT scan of a different patient, showing an obstructed concha bullosa on the left (arrowed).
patient complains of bouts of sneezing associated with watery rhinorrhea, nasal obstruction and sometimes conjunctival irritation. In the chronic form, nasal obstruction is the most prominent symptom, while discharge and sneezing are sometimes absent. Possible treatment includes allergen avoidance, specific immunotherapy, mast cell degranulation inhibitors, and antihistamine or steroid preparations.

Polyps appear as bilateral, glassy swellings that progressively fill the nose and are associated with chronic sinusitis, anosmia and chronic nasal discharge. The aetiology of nasal polyps is unknown, but they are not (as previously thought) due to atopy. In some patients, a dramatic response to systemic steroids may be maintained by the regular use of topical steroid drops. In others, the response is less satisfactory or more short-lived, and for these patients surgical removal of the polyps is required. However, polyps tend to recur, and it is not unusual for patients to have multiple surgical procedures throughout their lives. Nasal polyposis in a few patients is associated with hypersensitivity to salicylates, and their management may be improved with a special diet.

Rhinitis medicamentosa is an uncommon condition caused by the abuse of nasal vasoconstrictor medication. Some of these patients use 10 or more bottles of nasoconstrictor drops per day (Fig. N.13). Nasal obstruction is the predominant symptom, and it is extremely resistant to therapy. Topical steroid nasal drops can be effective, but care must be taken as they are absorbed extremely well by the chronically inflamed mucosa, and significant suprarenal suppression can result.

Vasomotor rhinitis or autonomic imbalance is a diagnosis of exclusion made when no other cause for nasal obstruction can be incriminated. The nasal vasculature undergoes cyclical and alternate changes (physiological nasal cycle). In this way, the mucosa of one or other nostril is congested while the other is relatively constricted. Vasomotor rhinitis is thought to be the derangement or exaggeration of this physiological process.

Wegener’s granulomatosis and mucous membrane pemphigoid may also affect the nasal mucosa where they present with gross crusting, spotting of nasal secretions with blood, impairment of the sense of smell and obstruction.

Nasal tumours, especially malignant neoplasms, are extremely rare. However, progressive unilateral nasal obstruction should always alert one to the possibility of a sinonasal tumour, particularly when associated with intermittent epistaxis and pain (Fig. N.14).
Tumours may develop in all age groups. A list of the neoplasms found in the nose is provided in Box N.2. The diagnosis is made on the basis of clinical examination and biopsy.

**NASAL REGURGITATION**

Michael Gleeson

Most people have experienced the regurgitation of food or drink through their nose at some time or other in their life. It is most unpleasant and is usually caused by a lapse of concentration when swallowing. Progressive and persistent nasal regurgitation may be caused by a number of conditions that are classified in Box N.3.

Chronic pathological nasal regurgitation occurs when either the soft palate is too short, too rigid or paralysed, or when there is a defect in the hard palate, alveolus or buccal sulcus.

Abnormalities of the soft palate preclude complete closure of the postnasal space during swallowing. A cleft palate is the most common structural cause, and this is almost always diagnosed directly after birth. Surgical correction is not always entirely successful, and minor degrees of regurgitation may persist. Subtle abnormalities of the soft palate, for example submucosal clefts, may not be diagnosed for some time. Removal of the adenoids in all of these patients can exacerbate regurgitation. Less common causes include post-tonsillectomy scarring and over-enthusiastic uvulopalatopharyngoplasty.

Oro-antral fistulae arising following difficult dental extractions when the floor of the axillary anthrum is damaged or fractured. The majority close spontaneously, but occasionally some persist and require a secondary surgical closure. Other reasons for structural defects of the palate are listed in Box N.3.

The motor nerve supply of the palate is through the pharyngeal plexus, of which the major contribution is from the vagus. The sensory supply is shared by the Vth and IXth nerves. Interruption or interference with any of these nerves may result in regurgitation. Damage to the vagus and glossopharyngeal nerves may be caused by lesions within the posterior fossa, jugular foramen or parapharyngeal space. Lesions responsible for damage to the maxillary division of the Vth nerve are usually found in the floor of the cavernous sinus or pterygopatine fossa.

Paralysis of the palate, when bilateral, may be difficult to notice at rest, but it remains immobile on phonation and is even more obvious if the patient is made to gag. In addition, the patient is unable to blow up a balloon. When the paralysis is unilateral, the normal side is drawn up like a curtain, and the uvula is displaced to the normal side.

**Box N.3 Causes of nasal regurgitation**

<table>
<thead>
<tr>
<th>Structural abnormalities of the palate</th>
<th>Paralysis of the soft palate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Congenital</td>
<td>Damage to the nucleus of the Xth cranial nerve</td>
</tr>
<tr>
<td>• Cleft palate</td>
<td>• Posteroinferior cerebellar artery thrombosis</td>
</tr>
<tr>
<td>• Surgical</td>
<td>• Tumours of the medulla</td>
</tr>
<tr>
<td>• Oro-antral fistula after dental extraction</td>
<td>• Bulbar palsy</td>
</tr>
<tr>
<td>• Palatal fenestration following surgery for malignancy of maxillary antrum</td>
<td>• Polymyelitis</td>
</tr>
<tr>
<td>• Non-surgical</td>
<td>• Landry’s ascending paralysis</td>
</tr>
<tr>
<td>• Attempted suicide</td>
<td><strong>Posterior fossa lesions</strong></td>
</tr>
<tr>
<td>Inflammatory</td>
<td>• Tumours</td>
</tr>
<tr>
<td>• Destruction or perforation of palate due to syphilis, tuberculosis or leprosy</td>
<td>• Syphilitic meningitis</td>
</tr>
<tr>
<td>• Mucormycosis</td>
<td>• Glomus jugulare tumour</td>
</tr>
<tr>
<td>Malignant disease</td>
<td>• Hydrocephalus</td>
</tr>
<tr>
<td>Fixation of the soft palate</td>
<td><strong>Extracranial lesions</strong></td>
</tr>
<tr>
<td>Post-surgical deformity</td>
<td>• Malignant disease</td>
</tr>
<tr>
<td></td>
<td><strong>Miscellaneous</strong></td>
</tr>
<tr>
<td></td>
<td>• Post-diphtheritic paralysis</td>
</tr>
<tr>
<td></td>
<td>• Myopathies</td>
</tr>
<tr>
<td></td>
<td>• Myasthenia gravis</td>
</tr>
</tbody>
</table>

**NAUSEA**

Simon Anderson

The term ‘nausea’ reflects a constellation of unpleasant symptoms including bloating, postprandial fullness, early satiety and a sensation of an impending need to vomit. Nausea may be acute or chronic and develops from a disorder in the gastrointestinal tract, a systemic/metabolic disease or the central nervous system. The majority of cases are trivial, the cause evident, and their resolution spontaneous.

In order to determine a cause of nausea, it is important to consider the potential physiological mechanisms, namely luminal (noxious stimulus), gut wall (mucosa, muscle and nerve endings), local feedback pathways (splanchnic nerves) and higher neurological centres.

The causes can be divided into acute or chronic. Acute causes are usually those that cause a noxious insult to the upper gastrointestinal tract. These include dietary factors (fatty or spicy/hot foods); toxins (alcohol), medications of any type, especially chemotherapy, non-steroidal anti-inflammatory...
NAUSEA

Drugs (NSAIDs), metronidazole, erythromycin and cholestyramine; acute gastroenteritis, including bacterial, parasitic (e.g. amoebiasis, giardiasis) or viral (Norwalk virus, rotavirus, norovirus, Epstein–Barr virus or HIV) causes. Agents that cause acute nausea without direct injury to the upper gastrointestinal tract include anaesthetic agents and non-prescribed drugs, particularly marijuana. Vestibular neuritis (or labyrinthitis) is a well-recognized non-gastrointestinal cause of acute nausea and often vomiting. The cyclical vomiting syndrome is characterized by acute stereotypical episodes of self-limiting nausea and vomiting.

Chronic nausea can also occur due to injury to the upper gastrointestinal tract especially peptic ulcer disease (gastritis, duodenitis, reflux esophagitis or peptic ulceration; *Helicobacter pylori*), infections (CMV, tuberculosis); medication (as above). Motility disorders including gastroparesis (due to type I diabetes or idiopathic) or chronic idiopathic intestinal pseudo-obstruction should be considered. Infiltrations including eosinophilic gastroenteritis and hyperplasia (Menetrier’s disease) are uncommon causes. Pregnancy and electrolyte disturbances (sodium, potassium) are common causes and easily overlooked.

Cancers which are almost invariably associated with other symptoms may also cause chronic nausea. These include gastric adenocarcinoma (especially if in the antrum and causing gastric-outflow obstruction), gastric mucosa-associated lymphoid tissue (MALT) lymphoma and pancreatic cancer.

Although surgery is usually associated with acute ileus, blind-loop or sump syndromes and their potential for associated bacterial overgrowth may underlie chronic nausea. A history of previous surgery (Billroth II gastrectomy, biliary surgery or gastrojejunostomy) is of significance. Some conditions present with pain and vomiting with nausea being a secondary symptom. These include mesenteric ischaemia (atherosclerosis, embolic, vasculitis) and mechanical obstruction (volvulus, intussusception, adhesions, strictures).

The gut has a limited repertoire of symptoms through which it can highlight pathology. A degree of malabsorption due to pancreatic exocrine insufficiency or coeliac disease/tropical sprue and the rare Whipple’s disease may result in nausea. The presence of inflammation in the bowel wall (Crohn’s disease, ulcerative colitis, ischaemic colitis and diverticulitis) can cause nausea on account of irritation of the visceral peritoneum.

Similarly, the presence of peritoneal deposits of tumour, endometriosis or the accumulation of ascites may also give rise to nausea. Metabolic disorders, whether endocrine (hyperparathyroidism, hyperthyroidism, pheochromocytoma), renal (acute/chronic renal impairment), hepatic (acute/chronic impairment) or manifesting as electrolyte imbalances of other causes (e.g. hypercalcaemia) commonly result in the onset of nausea.

Migraine is associated with recurrent nausea and headache. Other intracranial causes of nausea (hydrocephalus, tumour, infarct and middle-ear disease), although uncommon, must be considered in the absence of any other credible cause.

As the potential causes for nausea are myriad, a detailed history is vital to provide the starting point for further investigation. The history must detail setting (acute/chronic, during travel, exposure to toxins, relationship to meals, recent foreign travel, others with similar symptoms and possibility of pregnancy) and the presence of other gut symptoms (lower/upper bowel) or systemic features (flu-like illness, rash, joint pains, headache, general well-being). Past medical history (surgery, peptic ulcer disease, diabetes mellitus and previous malignancy) and medication history (some of which include NSAIDs, iron, azathioprine, sulphonamides, steroids, bisphosphonates, colestyramine, statins, potassium/vitamin supplements, chemotherapy, metronidazole, erythromycin, digoxin and hormones/ hormonal antagonists) are very important. A family history may provide some clue as to cause (coeliac disease, inflammatory bowel disease or muscular dystrophies).

The physical examination too must be thorough. In the absence of focal gastrointestinal symptoms, signs of systemic (metabolic and autoimmune) or vascular (atrial fibrillation) disease must be sought, and raised intracranial pressure must be excluded.

The broad nature of the conditions that give rise to nausea precludes a limited list of investigations. In the absence of further focal symptoms or signs, general tests such as urea/electrolytes, liver blood tests, serum calcium/phosphate, full blood count and erythrocyte sedimentation rate may provide some leads. Gastroscopy with biopsy will identify peptic ulcer disease, cancer, infiltrative conditions, viral inclusions and gastric stasis. A gastric emptying study using (ppm)Tc scintigraphy will detect gastroparesis. An ultrasound or CT scan of the abdomen and pelvis will identify focal pathology...
in the absence of focal signs. A CT brain may be needed if no other cause is found, to exclude space occupying lesions and raised intracranial pressure.

NECK, ENGORGED VEINS IN

Gerry Carr-White

For practical purposes, the jugular venous pressure may be regarded as raised if jugular pulsation is visible, or the internal jugular vein is seen to be distended, with the patient sitting erect. This is more reproducible than the older criterion of 4 cm above the sternal angle with the patient lying at 45°, and a grossly elevated venous pressure is less likely to be missed. The first major distinction is between non-pulsatile and pulsatile elevation of the venous pressure.

NON-PULSATILE ELEVATION OF THE JUGULAR VENOUS PRESSURE

This is due to obstruction of the jugular or brachiocephalic vein or, most commonly, of the superior vena cava (SVC). The causes of this are listed below:

- Common causes:
  - Thrombosis following implantation of a pacemaker/defibrillator
  - Bronchial neoplasm
  - Mediastinal tumour (e.g. lymphoma or thymoma)
- Less common causes:
  - Fibrosing mediastinitis
  - Right atrial or pericardial tumours
  - Late complication of Mustard’s operation for transposition of the great arteries

The diagnosis of venous obstruction can be confirmed by Doppler ultrasound or venography; the cause of the obstruction may be more difficult to diagnose, and needs further investigation with chest radiography supplemented by computed tomography or magnetic resonance imaging.

PULSATILE ELEVATION OF THE JUGULAR VENOUS PRESSURE

This condition is much more common, the causes including:

- Common causes:
  - Congestive cardiac failure
  - Overtransfusion
  - Cor pulmonale/right heart failure
  - Tricuspid regurgitation
- Less common causes:
  - Pericardial effusion
  - Massive pulmonary embolism
  - Constrictive pericarditis
  - Tricuspid stenosis
  - Restrictive cardiomyopathy
- Rare causes:
  - Carcinoid syndrome
  - Right atrial tumours

An analysis of the venous waveform helps to identify the cause of the elevation (Table N.1). A ‘normal’ jugular pulse has three ‘humps’ (the a, c and v waves) and two ‘dips’ (the x and y descents). The a wave is transmitted atrial systole, the c wave occurs early in systole as the tricuspid valve closes and braces back towards the atrium, and the v wave is produced by filling of the atrium from the venae cavae followed by emptying of the atrium after tricuspid valve opening (Fig. N.15). The relationship of this pattern to those found in disease is not always straightforward. Where an increased load on the right heart develops gradually – as in pulmonary/tricuspid stenosis or some forms of pulmonary hypertension – the right atrium has time to hypertrophy, and there may be a prominent a wave. If the tricuspid valve becomes incompetent, the jugular pulse shows a systolic pulsation in time with the arterial pulse. This is traditionally called a v wave, although its timing and cause are both different from the ‘normal’ v wave.

Table N.1 Abnormalities of the jugular venous pulse

<table>
<thead>
<tr>
<th>Cause</th>
<th>Most prominent feature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Congestive failure</td>
<td>Raised mean pressure with early diastolic descent</td>
</tr>
<tr>
<td>Tricuspid regurgitation</td>
<td>Systolic v waves</td>
</tr>
<tr>
<td>Right atrial hypertrophy</td>
<td>Presystolic a waves</td>
</tr>
<tr>
<td>Complete heart block, ventricular tachycardia</td>
<td>Cannon waves (sporadic or regular sharp v waves), pacemaker, etc.</td>
</tr>
<tr>
<td>Constrictive pericarditis</td>
<td>Very high pressure, sharp early diastolic descent</td>
</tr>
<tr>
<td>Tricuspid stenosis</td>
<td>High pressure, slow diastolic descent</td>
</tr>
</tbody>
</table>
The v waves of tricuspid regurgitation are often striking in appearance and readily palpable. Cannon waves are also prominent systolic waves appearing in the venous pulse, but these are due to contraction of the atrium against a closed tricuspid valve. Classic cannon waves are seen in patients with complete heart block and atrioventricular dissociation, when the atria and ventricles contract at their own independent frequencies. When atrial and ventricular contractions coincide, the cannon waves are seen as striking, sharp pulsations in the jugular pulse. Cannon waves may also occur in patients in junctional rhythm and in those with a permanent ventricular pacemaker who have preserved retrograde atrioventricular conduction; in such cases, retrograde atrial activation and cannon waves may occur in a series of consecutive beats. The murmur of tricuspid regurgitation is, of course, absent.

When the mean jugular venous pressure is considerably raised, the sudden fall in pressure (and consequent collapse of the vein) as the tricuspid valve opens at the beginning of ventricular diastole (y descent) is usually the most prominent feature of the venous pulse. In patients with tricuspid stenosis, this fall in pressure occurs more slowly as the right ventricle fills via the stenotic valve. In patients with constrictive pericarditis, the y descent is usually prominent, but a proportion of patients show a more pronounced dip in early systole (x descent), possibly because ventricular contraction causes a transiently negative intrapericardial pressure. This may also be seen in pericardial tamponade, although many cases simply show a prominent y descent. Patients with constrictive pericarditis or pericardial tamponade may also show a paradoxical elevation of venous pressure on inspiration (Kussmaul’s sign). The probable mechanism is due to a corresponding increase in right ventricular filling pressure in a patient unable to increase the right ventricular stroke volume.

**NECK, PAIN AND/OR STIFFNESS**

Fred Heatley and Jonathan Lucas

These two symptoms usually co-exist, or else stiffness will be ‘residual’ to a previous painful episode, for example a cervical disc prolapse. The exception is the congenital causes, which are usually painless and present with stiffness accompanied by deformity. The causes of neck stiffness and/or pain are listed in Box N.4.

**CONGENITAL CAUSES**

(See also SPINE, DEFORMITY OF, p. 627.)

Congenital muscular torticollis or wryneck is due to a contracture of the sternocleidomastoid muscle on one side, and it is generally considered to be the result of an injury during obstetric delivery, possibly ischaemic in nature. The muscle stands out as a tight band in the neck, and its contracture leads to a characteristic deformity. The head is pulled down towards the affected side, and the face and chin are tilted towards the opposite shoulder. The movements of the head are necessarily restricted owing to the shortening of the muscle, which in long-standing cases leads to a marked asymmetry of the face. The consequences are not limited to the head and neck, as the spine shares in the general obliquity and shows marked lateral curvature.
in old cases. This condition must not be confused with a secondary torticollis, which is an acquired condition and may indicate underlying disease, for example vertebral injury, pyogenic infection, intraspinal tumour, intracranial tumour of the posterior fossa, or unusual bone pathology (e.g. an osteoid osteoma).

In the Klippel–Feil syndrome there is a congenital fusion of one or more cervical vertebrae resulting in a short, thick, stiff neck, the head being set low on the shoulders. Other co-existing abnormalities are common: undescended scapulae (Sprengel’s deformity), platybasia, etc. (see UPPER LIMB, PAIN IN, Fig. U.3, p. 703).

**ACQUIRED ACUTE CAUSES**

**Traumatic causes**

Exposure to cold – for example, a cold draught from air-conditioning or sleeping in a cramped position – may give rise to a transient stiff neck, which is associated with no other symptoms. The patient wakes up in the morning with a stiff neck, and the diagnosis is made by exclusion. The symptoms settle rapidly.

Fractures and dislocations may be rapidly fatal, particularly when they affect the upper cervical spine. A classic example is a fracture through the pedicles of C2, often referred to as the ‘hangman injury’. Road traffic accidents are the most common cause. A second group of patients present with associated cord or nerve root compression, and consequently there is a clear pointer to the underlying injury. Beware, however, the third group, in which there may be major instability but no abnormal neurology and apparently normal plain X-rays. These injuries are easy to miss, but the patient is of course still at major risk of developing a paralysis should they suffer a simple fall. These are by no means all high-velocity injuries, but they may occur following relatively simple falls and landing on the forehead (e.g. falling down stairs, skiing, playing sports, etc.). A common sign in this third group is that the patient supports the head with their hands. All those patients who have significant trauma or are suspected of having significant head or neck trauma should have a lateral cervical spine x-ray as part of their initial assessment as per the ATLS guidelines. The cervicothoracic junctional zone is difficult to visualize in broad-shouldered men. Imaging with CT and MRI scans are very helpful in making a precise diagnosis.

Whiplash injuries have not only become a common cause of cervical pain, but are also a major source of medico-legal compensation. The majority are due to a rear impact collision to a car, causing a hyperextension injury of the neck. The precise soft-tissue pathology is a matter of dispute, but the disc, the anterior longitudinal ligament and the facet joints have all been implicated.

Neck pain and headaches are the most common symptoms and often appear disproportionately great in comparison to the physical signs. Most patients recover within 2–3 months. Some take 1–2 years to resolve, while a few are left with a permanent disability.

**Acute local infection**

Acute pyogenic infection of the cervical spine is uncommon but is increasing in incidence secondary to infection around intravenous cannulae. It is usually staphylococcal in nature. In the early stages, it is extremely easy to overlook and, as with other infections in the spine, it may present as a pyrexia of unknown origin.

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**Box N.4** Causes of neck stiffness and/or pain

<table>
<thead>
<tr>
<th>Congenital</th>
<th>Degenerative</th>
</tr>
</thead>
<tbody>
<tr>
<td>Congenital (infantile) torticollis or ‘wryneck’</td>
<td>Acute cervical disc prolapse</td>
</tr>
<tr>
<td>Congenital deformities (e.g. Klippel–Feil syndrome)</td>
<td>Acute painful episode in cervical spondylitis</td>
</tr>
<tr>
<td><strong>Acquired: acute</strong></td>
<td><strong>Miscellaneous</strong></td>
</tr>
<tr>
<td><strong>Traumatic</strong></td>
<td>Exposure to cold</td>
</tr>
<tr>
<td>Fractures, dislocation and subluxations of the cervical spine</td>
<td>Positional</td>
</tr>
<tr>
<td>Soft-tissue injuries to muscles and ligaments including whiplash injury</td>
<td>Intra-cervical and subarachnoid haemorrhage</td>
</tr>
<tr>
<td><strong>Infective: local</strong></td>
<td>Cerebral tumour</td>
</tr>
<tr>
<td>Acute pyogenic infection</td>
<td>Hysteria (remember, this is diagnosed by ‘excluding’ other causes)</td>
</tr>
<tr>
<td>Abscess in the neck</td>
<td><strong>Acquired: chronic</strong></td>
</tr>
<tr>
<td>Reflex spasm due to adenitis from otitis media, tonsillitis, etc.</td>
<td>Untreated acute traumatic lesions</td>
</tr>
<tr>
<td><strong>Infective: systemic</strong></td>
<td>Contractures following burns, nerve injuries, etc.</td>
</tr>
<tr>
<td>Meningitis</td>
<td><strong>Degenerative</strong></td>
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<tr>
<td>Typhus</td>
<td>Cervical spondylitis</td>
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<tr>
<td>Brain abscess</td>
<td>Ossification of the posterior longitudinal ligament</td>
</tr>
<tr>
<td>Poliomyelitis</td>
<td><strong>Arthritic</strong></td>
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<tr>
<td>Psittacosis</td>
<td>Chronic juvenile arthritis (Still’s disease)</td>
</tr>
<tr>
<td>Arbovirus infections (e.g. sandfly fever)</td>
<td>Rheumatoid arthritis</td>
</tr>
<tr>
<td>Leptospirosis</td>
<td>Ankylosing spondylitis</td>
</tr>
<tr>
<td>Tetanus, etc.</td>
<td>Other spondylarthropathies</td>
</tr>
<tr>
<td><strong>Malignant</strong></td>
<td><strong>Post-traumatic</strong></td>
</tr>
<tr>
<td>Metastatic neoplasms (N.B. primary neoplasms are exceptionally rare)</td>
<td>Untreated acute traumatic lesions</td>
</tr>
<tr>
<td>Multiple myeloma</td>
<td>Contractures following burns, nerve injuries, etc.</td>
</tr>
<tr>
<td><strong>Miscellaneous</strong></td>
<td><strong>Degenerative</strong></td>
</tr>
<tr>
<td>Cerebral tumour</td>
<td>Cervical spondylitis</td>
</tr>
<tr>
<td>Hysteria (remember, this is diagnosed by ‘excluding’ other causes)</td>
<td>Ossification of the posterior longitudinal ligament</td>
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<tr>
<td><strong>Acute traumatic</strong></td>
<td><strong>Arthritic</strong></td>
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<tr>
<td><strong>Infective</strong></td>
<td>Chronic juvenile arthritis (Still’s disease)</td>
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<tr>
<td><strong>Degenerative</strong></td>
<td>Ankylosing spondylitis</td>
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<tr>
<td><strong>Arbovirus infections</strong> (e.g. sandfly fever)</td>
<td><strong>Miscellaneous</strong></td>
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Inflammation around the cervical lymph nodes commonly causes reflex muscle spasm. The primary infection may be obvious – for example, a boil or carbuncle – but do not overlook carious teeth or infected tonsils.

Acute systemic infection
Many acute infections are accompanied by a stiff neck (meningism), particularly in children; pneumonia, once a common cause in childhood, is now much less so. Fever is almost always present. Meningitis from any cause, bacterial or viral, almost always causes some neck rigidity. Neck stiffness may be an early prodromal sign in paralytic or non-paralytic poliomyelitis, and changes in the cerebrospinal fluid are found. Phlebotomus (sandfly) fever – an arbovirus infection – presents as fever, malaise, myalgia and sometimes headache, in some cases with findings of an aseptic meningitis. Stiffness of the neck may be an early sign of tetanus, but other signs such as trismus (inability to open the mouth due to tonic contraction of the jaw muscles) rapidly appear (see TRISMS (LOCKJAW), p. 695).

Malignant causes
By far the most common group of tumours affecting the cervical spine are metastatic deposits. In order of frequency, these are carcinoma of the breast, prostate, lung, kidney and thyroid, but rarely of the bladder and gastrointestinal tract. By the time the patient is symptomatic, destructive or sclerotic X-ray changes are usually obvious, although sometimes it is easy to overlook the destruction of a pedicle (see SPINES, TENDERNESS OF, p. 634; Figs S.54 and S.55). By contrast, X-rays in multiple myeloma may show nothing more than osteopenia until a vertebral body collapses (Fig. N.16). Benign tumours of the cervical spine are uncommon, but are a differential diagnosis of a cervical disc presenting with radiculopathy or myelopathy. An example is a neurofibroma producing nerve root compression at an exit foramen.

Degenerative causes
An acute cervical disc is often precipitated by a local strain, especially sudden unguarded flexion and rotation. In essence, there are three clinical pictures:

- Symptoms and signs confined to the cervical spine
- Pressure on a nerve root with radicular symptoms radiating into an arm
- A central prolapse with pressure on the cord producing myelopathy (for further details, see UPPER LIMB, PAIN IN, Box U.2, p. 698)

Miscellaneous causes
Cerebral or subarachnoid haemorrhage
Conscious patients who have intracranial bleeding usually complain of headache and stiffness of the neck. Indeed, after subarachnoid haemorrhage, the main physical sign is marked neck rigidity and pain on trying to move the head. A brain tumour may cause a stiff neck due to meningeal irritation from bleeding into the subarachnoid space, by direct meningeal involvement or by causing cerebellar herniation through the foramen magnum.

Hysteria
A theatrical and over-dramatic symptom of neck stiffness is accompanied by other features of hysteria, but not by any objective physical signs of organic disease.

ACQUIRED CHRONIC CAUSES
Degenerative causes
Cervical spondylosis is the most common disorder of the cervical spine. The symptoms come on gradually over many years, and are typically episodic, acute

Figure N.16 Myelomatous destruction of C6: (a) lateral X-ray, and (b) magnetic resonance imaging scan.
exacerbations interspersed with long periods in which underlying stiffness only causes minor inconvenience. As with an acute disc, symptoms essentially fall into three patterns:

- Neck pain with radiation up into the occiput, out over the shoulder or down the thorax over the scapula
- Radicular pain secondary to osteophytic impingement and narrowing of an exit foramen
- Myelopathy due to central impingement on the cord, often from multilevel disease

This third group, which particularly affects the elderly, is often of insidious onset, and neurological features – particularly unsteadiness of gait – are all too easy to attribute to ‘old age’. Radiological changes in the neck are the norm in those aged over 60, so care must be taken in relating radiology to symptoms. If neurological compression – either of a nerve root or the cord – is suspected, an MRI scan is the investigation of choice.

Ossification of the posterior longitudinal ligament is a rare disease, with the majority of cases having been reported from Japan. In addition to neck pain and stiffness, nerve root compression is common, and myelopathy may result from central compression as the ossification increases.

Rheumatoid arthritis involves the cervical spine in as many as 30 per cent of cases. Rheumatoid erosion leads to instability, which can be a cause of sudden death. The myelopathy is not easy to diagnose as the neurological deficiency in the limbs is often masked by the peripheral joint disease. There are three categories: (i) basilar invagination with the odontoid (dens) migrating through the foramen magnum, affecting the brainstem and upper cervical cord; (ii) atlanto-axial instability usually with anterior displacement of C1 on C2 (Fig. N.17b); and (iii) subaxial instability, which can affect any part of the remainder of the cervical spine. Beware, therefore, of the patient with long-standing chronic rheumatoid arthritis (Figs N.17 and N.18) who presents with increasing pain in the cervical spine, a progression of the pain up over the occiput indicative of compression of the posterior primary rami, and symptoms such as tinnitus, vertigo, visual disturbances or dysphagia and l’Hermitte’s syndrome with flashing electrical shock-like pains involving all or part of the body and being indicative of cord compression.

Ankylosing spondylitis can extend up to involve the cervical spine (see BACK, PAIN IN, p. 51). Even when it becomes ‘burnt out’, the spine remains at increased risk of injury due to the loss of its flexibility. Do not therefore dismiss the recurrence of cervical spine pain in a patient with old ankylosing spondylitis as being due to a recurrence of the disease, as there may well be a non-united fracture that has occurred secondary to a relatively minor injury (Fig. N.19).

Cervical vertebral tuberculosis

The greatest care must be taken not to overlook tuberculous disease of the cervical vertebrae as a cause of reflex muscular rigidity of the neck. Pain and rigidity are among the earliest signs; the pain is increased by the least movement, and the child – as it is generally a child that is affected – takes the greatest precaution to avoid any movement, even holding the head between his or her two hands. The position of the head varies; it is most often held very stiff and straight, the natural backward curve of the neck being lost. In the late stages, there may be an angular or lateral curve.
NECK, SWELLING OF

Harold Ellis

(See also THYROID ENLARGEMENT, p. 678.)

ANATOMY

The neck on either side is divided into anterior and posterior triangles by the sternocleidomastoid muscle arising from the sternum, sternoclavicular junction and medial third of the clavicle below, and being inserted into the mastoid process of the temporal bone above. At the upper end of the anterior triangle, the digastric muscle defines the lower borders of a subsidiary space known as the ‘digastric triangle’, and at the lower end of the posterior triangle, the posterior belly of the omohyoid muscle defines the upper border of a subsidiary space known as the ‘supraclavicular fossa’.

The sternocleidomastoid muscles are enclosed within the deep cervical fascia, which splits to embrace them. If even a part of a mass in the neck overlaps either border of the sternocleidomastoid muscle, then, by putting one or other of these muscles into contraction, the relationship of the mass to the sternocleidomastoid muscle and so to the deep fascia can readily be determined. This method is applicable to practically all masses in the neck, except for the majority of those situated in the midline. The right sternocleidomastoid muscle is put into contraction by rotating the head to the left while resistance is applied to the chin, and vice versa; both sternocleidomastoids are made to contract when the forehead is pressed forwards against resistance.

Lumps in the neck arising from structures superficial to the deep cervical fascia are not specific to the neck. Thus, sebaceous cysts, lipomas, carbuncles, etc., are common, particularly in or deep to the skin at the back of the neck. It is the masses deep to the deep cervical fascia which have particular relevance in regard to the neck, and it is the differential diagnosis of these that must be considered.

It is conventional to divide swellings in the neck into midline swellings and lateral swellings, although this is a little misleading as nearly all so-called ‘midline swellings’ can deviate slightly to one side or the other. They can, however, be divided appropriately into masses arising from unpaired midline structures and masses arising from paired lateral structures.

MASSES ARISING FROM UNPAIRED MIDLINE STRUCTURES

Thyroglossal cyst

The thyroid gland is developed from an epithelium-lined duct that grows downwards from the region of the foramen caecum of the tongue, passing close in front of and then behind the hyoid bone, and so towards the site of the adult thyroid isthmus, from which the lateral lobes expand. A cyst may form in any part of this track by failure of obliteration of the duct, but the most common site is at the lower border of the hyoid bone, anterior to the thyrohyoid membrane. These cysts usually appear at about puberty and enlarge to a variable size
slightly to one or other side of the midline. They are fluctuant, globular masses that, if superficial, may transilluminate. If the jaw is held open and the tongue steadily protruded, the swelling will rise in the neck, demonstrating its attachment to the base of the tongue (Fig. N.20). These cysts occasionally become infected and may rupture, leading to a fistula.

Swellings arising from the isthmus of the thyroid gland
All of those pathological conditions described (see below) and which give rise to swellings of the thyroid gland can arise in the isthmus. It should be repeated that practically all thyroid swellings move up and down on deglutition owing to their intimate relationship to the larynx and upper part of the trachea, the movements of which they follow during this act.

Rare causes of swelling arising in midline structures
These include:
• Subhyoid bursa – a cystic swelling that arises behind the hyoid bone and is clinically indistinguishable from a thyroglossal cyst
• Perichondritis of the thyroid cartilage
• An advanced carcinoma of the larynx, trachea or oesophagus penetrating the walls of these viscera and protruding to one or other side
• The so-called ‘Delphic lymph node’, which lies in the midline on the thyrohyoid membrane; this may enlarge in carcinoma of the thyroid gland and may be the first evidence of this disease

MASSES ARISING FROM PAIRED LATERAL STRUCTURES

Lymph nodes
(See also LYMPHADENOPATHY, p. 394.)
The most common swellings in the neck are undoubtedly due to pathological processes arising in the lymph nodes, usually secondary to some acute or chronic inflammatory or a neoplastic process in one of the organs that they drain, but sometimes (as in the lymphomas) appearing to arise primarily within these nodes.
The distribution of the lymph nodes in the neck is variable, but the general disposition is as follows. In the upper part of the neck, there is a horizontally disposed system consisting of the submental, suprathyroid, submaxillary and upper deep cervical groups. The names of these groups indicate sufficiently
NECK, SWELLING OF

their situation, except for the upper deep cervical group, which is situated in relation to the internal jugular vein, where it is crossed by the posterior belly of the digastric muscle. One important node of this group – the jugulodigastric node – is particularly significant in relation to pathological conditions of the tongue and tonsil. As a general rule, the anterior nodes drain anteriorly placed structures in the head, while the posterior nodes drain more posteriorly situated organs. Thus, the tip of the tongue drains to the submental nodes, while the posterior aspect of the scalp drains to the occipital nodes.

In addition to the horizontal system, there is a vertical system ranged along the internal jugular vein. At the upper end, there is the upper deep cervical group, which is common to both systems, and at the lower end the lower deep cervical group with subsidiary groups in between. The lower deep cervical group of lymph nodes lies in relation to the internal jugular vein, where it is crossed by the posterior belly of the omohyoid, and one large node of this group – the jugulo-omohyoid node – is again of significance in relation to pathological processes in the tongue, receiving lymphatics from this organ without the interposition of any intervening lymphatic nodes. In this way, a carcinoma of the side of the tongue can give rise to secondary deposits in the supraclavicular fossa, where this node is situated, without the enlargement of any of the systems in the upper part of the neck. The Delphic node on the thyrohyoid membrane, which has already been referred to, should also be mentioned at this point. For a description of the differential diagnosis of the various types of enlarged lymph nodes, see page 394.

Thyroid swellings

These are the second most common cause of swellings situated laterally in the neck (see list on the right). Nearly all these swellings move up and down with deglutition, by which property they may be recognized. There are, however, some exceptions to this rule. If the mass is very large and fills one or both anterior triangles and perhaps also the midline, there may not be room for the thyroid to move on deglutition. Again, in certain types of carcinoma with infiltration of the pretracheal muscles, the growth may not move on deglutition. This is because the larynx, which causes the thyroid to move, cannot itself do so as there is no elasticity left in the infiltrated pretracheal muscles. Indeed, it is this tethering of the larynx by infiltration of the surrounding structures which leads to dysphagia in carcinoma of the thyroid, as can readily be appreciated by anyone who attempts to swallow while holding down their thyroid cartilage by placing a finger on its upper border. Sometimes, in a nodular goitre, the excursion of the mass on swallowing or on coughing may be so considerable that it rises up from ‘plunging down’ into the superior mediastinum or retroclavicular spaces during these movements. This so-called ‘plunging goitre’ is only a type of retrosternal or retroclavicular goitre with an abnormally free range of movement. Its very mobility argues that it will probably be a simple matter to deal with surgically.

More rare causes of swelling laterally placed in the neck

These may be listed as:

- Branchial cyst
- Sternomastoid tumour
- Cervical rib
- Cystic hygroma
- Aneurysm
- Carotid body tumour
- Submandibular salivary swellings (sialectasia or tumour)
- Actinomycosis
- Cold abscess from spinal tuberculosis
- Pharyngeal pouch
- Laryngcele

Branchial cyst

This is a congenital condition believed to arise in the remains of the second branchial cleft and to give rise to a cystic swelling in the lateral part of the neck. Another theory is that it represents cystic degeneration within a lymph node. The condition may arise at any age, but usually occurs in young people and is rare after the age of 40 (Fig. N.21a and b). The swelling, which varies in size, usually protrudes into the anterior triangle from the deep surface of the upper part of the sternocleidomastoid muscle. It is usually rather soft and fluctuates readily, but it is generally too deeply situated to demonstrate translucency or transillumination. Occasionally, these cysts become infected, when the differential diagnosis from breaking-down tuberculous nodes may be difficult. However, the diagnosis can usually be determined by aspiration, which will yield either tuberculous pus in the latter case or, on the other hand, turbid yellow fluid containing numerous cholesterol crystals on microscopic examination, which is typical of a branchial cyst (Fig. N.21c and d).

Although not strictly a swelling in the neck, it should be mentioned here that the unobliterated second branchial cleft, instead of forming a cyst, may communicate with
the exterior, usually just medial to the sternal head of the sternocleidomastoid muscle below and into the pharynx in the supratonsillar fossa above, forming a **branchial fistula**.

**Sternomastoid tumour**
As a result of birth or intrauterine injury, some fibres of the sternocleidomastoid muscle may be torn, and a haematoma appears in this muscle, leading to **torticollis** (wryneck).

**Cervical rib**
This is another congenital abnormality that may give rise to a swelling in the supraclavicular fossa. The swelling may be due to the rib itself, or there may be a pulsatile swelling due to a ‘post-stenotic’ dilatation of the subclavian artery (see p. 65 and Fig. B.8).

**Cystic hygroma**
This rare congenital abnormality is a lymphangiomatous condition that usually arises in the supraclavicular fossa of infants. It forms a soft, fluctuating, extremely translucent and painless swelling that may grow rapidly. These masses are liable to attacks of infection.

**Aneurysm and arteriovenous fistula**
The large vessels of the neck are liable to the same pathological changes as vessels elsewhere. Aneurysms may occur in the cervical part of the subclavian artery, or in the carotid arteries. A penetrating injury of the neck, as by a metallic fragment, may damage both the carotid artery and the internal jugular vein, leading to an arteriovenous fistula.

**Carotid body tumour**
This is a rare lesion arising in the chromaffin tissue situated at the bifurcation of the common carotid artery. It appears at any time after infancy as a very firm ‘potato-like’ tumour in close association with the carotid sheath, so that pulsation is usually, but

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**Figure N.21** Branchial cyst: (a) anterior view, and (b) lateral aspect. (c) Fluid aspirated from the cyst. (d) Cholesterol crystals seen on microscopic examination of this fluid.
NECK, SWELLING OF

Carotid body tumour: (a) anterior view, and (b) lateral aspect. (c) Carotid angiogram of a carotid body tumour to demonstrate the typical splaying apart of the external and internal carotid arteries. [Prof. Gerald Westbury.]

not invariably, transmitted to it (Fig. N.22). Its steady growth over a period of years serves to distinguish it from tuberculous cervical adenitis, with which it may readily be confused. Carotid angiography demonstrates the diagnostic splaying apart of the internal and external carotid arteries at their origins by the tumour mass at the bifurcation.

Swellings of the submandibular salivary gland
These arise in the digastric triangle (see SALIVARY GLANDS, SWELLING OF, p. 594). Ludwig’s angina is an acute inflammatory process of the cellular tissue around the submandibular gland, usually arising from the floor of the mouth or the teeth. The physical signs extend into the floor of the mouth and give rise to considerable oedema, which, without treatment, may spread to the glottis and demand tracheotomy.

Actinomycosis
This is a chronic inflammatory swelling of the cellular tissue about the angle of the mandible. The diffuse induration with the eventual development of multiple sinuses and the accompanying trismus should make the diagnosis obvious.

Late in the disease, ‘sulphur granules’ containing the streptothrix may be discharged, but the diagnosis should not await bacteriological confirmation, which may be equivocal in the early stages.

Cold abscess from spinal tuberculosis
In certain cases of tuberculosis of the cervical spine, the abscess may track from the retropharyngeal region laterally and present as a fluctuant mass in the upper part of the posterior triangle and deep to the insertion of the sternocleidomastoid muscle. The accompanying stiffness of the neck, together with the general evidence of a chronic infection, should alert the examiner to this possibility. If untreated, the abscess breaks down, discharges, and forms multiple sinuses in the apex of the posterior triangle. An X-ray examination will reveal typical destruction of the involved vertebra or vertebrae and adjacent disc.

Pharyngeal pouch
At the back of the inferior constrictor muscle of the pharynx, there is a triangular area (Killian’s dehiscence), located between the upper border of the transversely running fibres of the cricopharyngeus below and the lower border of the obliquely running fibres of the thyropharyngeus above, where the wall is deficient in muscle. A pouch of mucosa, covered only by the fascia propria of the pharynx, may protrude through this defect. This pouch gradually enlarges, usually towards the left side of the neck, and tends to fill up when food or fluid is swallowed. At first, this is just a nuisance, and gives rise to an uncomfortable feeling on swallowing together with a rapidly developing swelling, which may be emptied by pressing on the mass. This may be accompanied by regurgitation of fluid, often foul-smelling, into the pharynx and mouth. Later, the mass becomes sufficiently large to press upon the oesophagus, against which it lies, to produce severe dysphagia with inanition.
Food is apt to stagnate within the pouch, leading to diverticulitis that may spread and give rise to pharyngitis or oesophagitis, so adding to the burden of dysphagia. This condition may appear at any age, but it usually arises during the third and fourth decades of life. It is readily demonstrated by a barium swallow X-ray (Fig. N.23). Treatment, after attention to the nutritional needs of the patient, is by surgical excision.

Laryngocele
This is a narrow-necked, air-containing diverticulum consequent on herniation of the laryngeal mucosa through a defect of the thyrohyoid membrane, where this is pierced by the superior laryngeal vessels and the deep branch of the superior laryngeal nerve. Occasionally it is bilateral.

It appears as a swelling in the anterior triangle of the neck just above the lamina of the thyroid cartilage when the patient blows his or her nose – it may then remain distended for some days. It is likely to be found in professional trumpet-players and glass-blowers.

A lateral X-ray of the neck reveals the air-filled mass, also visible on computed tomography (Fig. N.24).

Deformities of the nipple may be classified as follows:

- **Congenital**
  - Congenital absence
  - Supernumerary
  - Congenital inversion
  - Bifid nipple (Fig. N.25)

- **Acquired**
  - Acquired inversion
  - Plasma cell mastitis
  - Mammillary duct fistula
  - Duct ectasia
  - Tumour

CONGENITAL ABNORMALITIES

Rarely, there can be complete absence of the development of the breast and the nipple. This may be associated with failure of development of the pectoral muscles, when the condition is known as Poland’s syndrome. Supernumerary nipple areolar complexes are quite common, running down the milk line from the subclavicular area across the lateral part of the abdomen, ending in the region of the anterior superior
iliac spine. Congenital inversions of the nipples are common, and must be distinguished from the acquired inversion of the nipple associated with either carcinoma of the breast or duct ectasia. A bifid nipple is a rare, but well-recognized, entity.

Acquired inversion of the nipple may be a consequence of duct ectasia/plasma cell mastitis syndrome (see BREAST LUMPS, p. 70). In this condition, inversion of the nipple is usually bilateral, central and slit-like. Apart from this, acquired inversion of the nipple is usually of sinister significance and may represent a retro-areolar carcinoma, or even the first sign of a cancer in one of the outer quadrants of the breast.

Paget's disease of the nipple is a relatively rare presenting sign of carcinoma of the breast and may be an eczematous condition affecting the nipple and areola (Fig. N.26). This is usually associated with an intraductal element of carcinoma invading along the terminal portions of the lactiferous ducts to infiltrate the dermis of the nipple and areola. If this eczematous condition is unilateral and not associated with patches of eczema elsewhere on the body, it should be treated seriously by an immediate biopsy. The histological appearance is characteristic, with foamy cells with large atypical nuclei seen scattered throughout the dermis and subdermal layers.

Mammillary duct fistula is a sequel of periductal mastitis, and presents as a discharging sinus at the areolar margin. For a description of this complex of diseases, see BREAST LUMPS, (p. 69).

Discharge from the nipple may be divided into three classes.

NORMAL DISCHARGES
A discharge of milk from the breast during pregnancy is not uncommon, especially in multiparae. Both then and during lactation, it is usually of small amount, except when the child is put to the breast, but occasionally the flow at other times may be sufficient to be distressing.

NORMAL DISCHARGES AT ABNORMAL TIMES
A secretion similar to colostrum sometimes occurs from the breasts of both sexes in the newly born, and again at puberty. This is due to endocrine stimulation, but it may predispose to a true infective mastitis when the breast, already tender and swollen, becomes hot and red, and the discharge may change from being clear to purulent.

Occasionally, the normal secretion of milk during lactation is prolonged for many months or years after the stimulus of suckling has been removed. This is probably due to some endocrine abnormality. Apart from being a serious nuisance, and sometimes also a source of anxiety to the patient, it has no sinister significance. It usually resolves spontaneously and
unpredictably after a varying period, with or without
the aid of endocrine therapy. Women with prolactin-
secreting tumours of the anterior pituitary may present
with galactorrhoea and amenorrhoea.

**ABNORMAL DISCHARGES**

**Serous fluid**
A discharge of serous fluid from the nipple is a
common accompaniment of duct ectasia, epithelial
hyperplasia or duct papilloma.

**Pigmented fluids**

**Green fluid**
When the colour is due to melanin or pigments other
than derivatives of haemoglobin, its admixture with
yellow serum gives to the resultant discharges a green
colour of varying shades. If the discharge is very dark,
dilution with water will disclose the green colour. In
cases of real difficulty, the discharge may be submitted
to spectroscopic or chemical assay for haemoglobin.
Such discharges have precisely the same significance
as the non-pigmented serous discharges discussed
above.

**Haemorrhagic**
Blood-stained discharges can usually be recognized
on sight; the colour is red to black, and again, if there
is real doubt, the final arbiters are the microscope
and the chemical test. Blood-stained discharges are
indicative of duct papilloma, epithelial proliferation and
intraductal carcinoma, in that order of frequency.

The nipple should be examined through a magnifying
glass, and a bead of blood or a speck of clot may reveal
from which of the 20 or so ducts the bleeding is arising.
Such evidence is important in determining from which
section of the breast the bleeding is originating. Having
examined the nipple thus, it should be wiped clean
and (with the breast rendered moderately tense by an
assistant if available) the tip of the finger pressed on
to the breast at successive sites, working spirally from
the nipple. Particular attention should be paid to the
subareolar region, where the source of the bleeding
lies in the majority of cases. By this means, it will be
found possible to cause blood to issue from the nipple
on pressure over quite a restricted area, whereas
pressure elsewhere has no effect. If the affected duct
has been previously identified, the significant area will
be found to be in the segment of the breast drained by
that duct, and the pathological region is confirmed. The
segment of the breast affected should be removed by
local operation, and the pathological condition causing
the bleeding determined by naked-eye inspection and
histological study. Further treatment depends upon
the nature of the lesion so determined. Solitary papillomas
adjacent to the nipple are the most common cause
of this symptom; if they are removed in this way, the
bleeding seldom recurs.

Should it be impossible to localize the origin of the
bleeding – and with care and practice this is most
unusual – the diagnosis depends on an assessment of
probabilities. The younger the patient, the more likely is
the cause to be benign; the older the patient, the more
likely to be malignant. Mammography is valuable in
demonstrating or excluding an occult neoplasm as the
source of the haemorrhage. Where the discharge of
pigmented fluids from multiple ducts associated with
duct ectasia is profuse and embarrassing, total excision
of the subareolar duct system (Hadfield’s operation)
will effect a cure.

**Grumous material**
The discharge of ‘cheese-like’ material or material
having the consistency of toothpaste or putty, grey
or green in colour, indicates the condition known as
‘comedo mastitis’. This is another variant of the duct
ectasia/periductal mastitis complex (see p. 70).

**Pus**
Pus, or pus mixed with milk, generally indicates acute
suppurative mastitis; the other signs of inflammation
or abscess are well marked as a rule, so that there is no
difficulty in arriving at a diagnosis. A tuberculous lesion
also causes a discharge of pus, and it may simulate
carcinoma. The discharge may contain demonstrable
tubercle bacilli, but specific bacteriological culture –
together with a radiograph of the chest – will very likely
be required before a positive answer on the nature of the
infection can be given.

**NODULES**

Barry Monk

A nodule is a solid palpable lesion in the skin, larger
than a papule (conventionally greater than 5 mm in
diameter). A large nodule may be called a tumour.
Nodules must be distinguished from cysts, which have
a fluid or semi-fluid content. The causes of nodular
skin lesions are many (Box N.5). Persistent non-tender
nodules should always be biopsied.

**NEOPLASMS**

Most nodular skin neoplasms are aetiologically
related to ultraviolet light (sunlight) exposure and
are thus most common on exposed areas of skin, in
fair-skinned subjects, and especially in those who
already show evidence of other ultraviolet-induced
changes in the skin. The most common tumour is a
basal cell carcinoma. Although these are also known
as rodent ulcers, ulceration is not invariably a feature
and is often a relatively late stage. Lesions begin as a small, rounded, pearly, translucent papule, showing telangiectasia. The head and neck are the most common sites, and the incidence rises with increasing age; lesions occurring below the age of 40 years are uncommon, although not unknown. Nodulocystic lesions grow as solid tumours composed of lobulated masses of cells in which cystic degeneration may occur. Enlarging lesions eventually ulcerate centrally, where a haemorrhagic crust forms. The morphoeic variant of rodent ulcer may erode widely, with a poorly defined infiltrating margin, and may thus be difficult to eradicate surgically.

A squamous cell carcinoma is a harder, fleshy nodule, often with overlying crust, generally arising on the face, including the lips and ears, or the dorsa of the hands (curiously an uncommon site for basal cell lesions). They tend to grow in a more aggressive fashion, to become secondarily infected (often with an unpleasant odour), and to metastasize to draining lymph nodes. They are especially common in patients who are immunosuppressed (e.g. following transplant surgery). Occasionally, squamous cell carcinoma arises in an old scar or sinus, or on the margin of a chronic leg ulcer; in such a case, it is easy to overlook the true diagnosis unless one is especially vigilant.

A kerato-acanthoma is now thought to be a variant of squamous cell carcinoma, but it has an unusual natural history. This lesion presents acutely as a neat, dome-shaped nodule that grows alarmingly in weeks on the exposed skin of the face and limbs.

A juicy hyperkeratotic central plug forms which may discharge later, giving the lesion the appearance of a giant molluscum contagiosum. Although some cases of kerato-acanthoma resolve spontaneously leaving a small irregular scar, some follow a more aggressive cause, so that they should be managed as with an invasive squamous cell carcinoma.

Nodular malignant melanomas are usually ominously obvious, although the amelanotic variant is often first diagnosed by the histopathologist (Figs N.27 and N.28). Early changes in moles that can point to malignant

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**Box N.5 Causes of nodular skin lesions**

<table>
<thead>
<tr>
<th>Neoplasms</th>
<th>Metabolic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Basal cell carcinoma</td>
<td>Xanthoma</td>
</tr>
<tr>
<td>Squamous cell carcinoma</td>
<td>Gouty tophus</td>
</tr>
<tr>
<td>Keratoacanthoma</td>
<td>Lipid proteinosis</td>
</tr>
<tr>
<td>Malignant melanoma</td>
<td>Pretribial myxoedema</td>
</tr>
<tr>
<td>Lymphoma</td>
<td></td>
</tr>
<tr>
<td>Kaposi’s sarcoma</td>
<td></td>
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<tr>
<td>Dermatofibroma</td>
<td></td>
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<tr>
<td>Pyogenic granuloma</td>
<td></td>
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<tr>
<td>Keloid</td>
<td></td>
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<tr>
<td>Vasculitis</td>
<td></td>
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<tr>
<td>Erythema nodosum</td>
<td>Mycobacteria</td>
</tr>
<tr>
<td>Nodular vasculitis</td>
<td>Fungi</td>
</tr>
<tr>
<td>Wegener’s granulomatosis</td>
<td>Legrosy</td>
</tr>
<tr>
<td>Polyarteritis nodosa</td>
<td>Treponema</td>
</tr>
<tr>
<td>Temporal arteritis</td>
<td>Oncocerciasis</td>
</tr>
<tr>
<td>Chronic inflammation</td>
<td>Infestation (including post-scabetic nodules)</td>
</tr>
<tr>
<td>Sarcoid</td>
<td></td>
</tr>
<tr>
<td>Rheumatoid nodules</td>
<td></td>
</tr>
</tbody>
</table>

**Infections**

- Acne
- Nodular prurigo

**Miscellaneous**

- Acne
- Nodular prurigo
- Keratoacanthoma
- Squamous cell carcinoma
- Keratoacanthoma
- Malignant melanoma
- Lymphoma
- Kaposi’s sarcoma
- Dermatofibroma
- Pyogenic granuloma
- Keloid
- Vasculitis
- Erythema nodosum
- Nodular vasculitis
- Wegener’s granulomatosis
- Polyarteritis nodosa
- Temporal arteritis
- Chronic inflammation
- Sarcoid
- Rheumatoid nodules

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Figure N.27 Malignant melanoma with cutaneous secondaries.

Figure N.28 Primary malignant melanoma.
change include haemorrhage, loss of hairs growing from the mole, pigment spilling into surrounding skin, ulceration and inflammation. *Lymphomatous infiltrates* in the skin are often nodular, and the later stages of *cutaneous cell lymphoma* (mycosis fungoides) are nodular and tumorous. *Kaposi’s sarcoma* comprises slow-growing, port-wine-coloured plaques and nodules, particularly on the lower limbs. A more aggressive form occurs widely on the body in HIV-positive individuals (and is now an accepted AIDS-defining diagnosis). Secondary deposits of an internal malignancy may present in the skin as hard dermal nodules (Fig. N.29); the diagnosis may be confirmed histologically, but it is usually a late presentation of widespread metastatic disease.

A common benign nodular neoplasm is a *dermatofibroma* (histiocytoma), which develops as a firm red/brown nodule in the upper dermis fixed to overlying skin usually on the legs (Fig. N.30). Histology shows a dense proliferation of fibrocytes, which may represent a tissue reaction to a preceding insect bite. Multiple lesions are sometimes encountered.

A *pyogenic granuloma* is a firm, small, cherry-red pedunculated nodule of hypertrophic granulation tissue, which bleeds easily on slight trauma and occurs most frequently on the lips and extremities. Its rapid growth and characteristic collarette are helpful diagnostic pointers, but histological confirmation is essential.

**VASCULITIS**

A cutaneous vasculitis commonly presents as multiple non-blanching vascular lesions in the skin; there may be associated features of systemic vasculitis. *Erythema nodosum* occurs most often in females in their second and third decades of life as crops of painful, tender, red nodules. These occur on the shins, are 1–8 cm in diameter, and heal over several weeks without breaking the surface and going through the colour changes of a bruise. A cause for this reaction can be found in half of the affected individuals. Examples include sarcoidosis, inflammatory bowel disease, infections (*Streptococcus*, tuberculosis, leprosy and deep fungus) and drugs (sulphonamides and thiazides).

*Nodular vasculitis* is also more common in women, but occurs later in life, in their third and fourth decades. The calves are the usual site of well-demarcated, bluish, fixed, subcutaneous nodules. Underlying tuberculosis must be excluded, but often a cause is not determined. *Polyarteritis nodosa* affects adult men, with nodules usually occurring along the course of arteries. There is severe illness with fever, arthralgia, hypertension, peripheral neuropathy and eosinophilia. The prognosis is related to the degree of renal vascular involvement. In *giant-cell arteritis*, exquisitely tender, nodular swellings occur most commonly along the course of the temporal artery. Sometimes, extensive ischaemic scalp ulceration is seen. Early recognition and treatment is important, as irreversible retinal artery thrombosis – and consequent blindness – can occur.

**CHRONIC INFLAMMATION**

Non-infectious granulomatous nodules are seen in sarcoidosis, rheumatoid arthritis, and around ruptured pilosebaceous glands in acne. *Sarcoidosis* usually begins in early adult life, more commonly in women, and with marked racial and geographical variation in prevalence. Skin presentations are varied, but they include dermal nodules, papules and infiltration of scars. *Pyoderma faciale* (Fig. N.31) is an aggressive variant of rosacea with tender nodules. *Erythema nodosum* is a common presentation of acute sarcoid (Fig. N.32). The nodules of *rheumatoid arthritis* occur over the bony prominences at points of pressure, such
as on the hands (Fig. N.33) and especially just below the elbow. They are painful, very rarely ulcerate, and vary in size from a pinhead to 2 cm in diameter. When present, they indicate seropositivity (Rose–Waaler or latex). Reaction to ingested bromides and iodides (bromoderma or iododerma) can rarely produce dramatic purplish, nodular and vegetative lesions on the face of infants or extremities of adults. The lesions may persist for long periods until recognized; they are now a rarity with the decline in use of these agents.

Firm, suppurating nodules are seen in nodulocystic acne vulgaris, due to granulomatous reactions around ruptured, swollen sebaceous glands. Similar lesions in the axillae, groins and perianal skin are seen in hidradenitis suppurativa, but these abscesses are based on apocrine glands. Young adults are chiefly affected, and the nodules are accompanied by draining sinuses.

**METABOLIC**

The deposition of metabolic products in the dermis may cause nodular swelling. These begin as papules, and include xanthoma, pretibial myxoedema, gouty tophi and lipoid proteinosis. Xanthomas are reddish-brown nodules of varying size, and are usually found on the elbows and knees and dorsa of hands and feet. They indicate an underlying disturbance of carbohydrate/lipid metabolism, usually primary but sometimes secondary (e.g. uncontrolled diabetes).

Gouty tophi are hard, yellow–white waxy nodules, on the helix or antihelix of the ear, palms or soles, tarsal plates of the eyelids and tendons of the hands or feet (Fig. N.34). Sometimes, tophi ulcerate and discharge a ‘cheesy’ material containing crystals of sodium biurate.

Lipoid proteinosis is a very rare inherited tendency to infiltrate the skin and mucosa with a hyaline material (mucopolysaccharide). Affected patients have a hoarse voice and characteristic beading along the eyelid margins.

In pretibial myxoedema, firm red nodules or plaques arise on the lower legs, over the front of the shin, and on the dorsa of the feet. The hair and follicle orifices are grossly hypertrophied, giving a peau d’orange appearance.
INFECTIONS

Nodules due to infection are less common in Western countries, except for the ubiquitous furuncle or boil. Boils appear chiefly on the face, neck and buttocks, and are painful red nodules, often with a yellow pustule at the apex. The cause is a follicular infection with Staphylococcus aureus, and lesions may be recurrent for many weeks or months. They are more frequent in diabetic and debilitated people.

Any organism which induces a chronic granulomatous tissue reaction seems particularly prone to produce skin nodules (see Box N.5).

Mycobacteria

Lupus vulgaris is now extremely uncommon. Small, softish, reddish-brown or yellowish nodules appear on the face or mucous membrane in childhood. When examined compressed under a glass slide (diascopy), the nodules have a typical ‘apple-jelly’ appearance. The nodules progress and slowly coalesce over very many years to form annular scaling plaques with central atrophy and fibrosis. More common now are nodules caused by infections with atypical mycobacteria, for example fish-tank granuloma (Fig. N.35). Usually, a finger is abraded against an infected fish-tank, and soft subcutaneous nodulocystic lesions develop. These are followed by a succession of nodules appearing along a lymphatic chain (sporotrichoid spread).

Fungi

In the UK, fungi and other infections in the immunosuppressed – particularly those with HIV disease – should be thought of as a cause of nodules. Actinomycosis causes abscesses around carious teeth, on the tonsils or in the gastrointestinal tract. The pus contains ‘sulphur’ granules of the penicillin-sensitive ‘ray fungus’, Actinomyces israelii. Cryptococcosis (torulosis) occurs in all parts of the world, usually involving the central nervous system, where meningitis, abscess or brain tumour may be suggested. Sometimes, transient skin nodules occur. Sporotrichosis begins with a skin nodule at the site of injury of hand or forearm, in contact with infected wood, soil or plant. There may be a considerable local inflammation before the characteristic succession of nodules appears up the draining lymphatic chain. Occasionally, a superficial dermatophyte infection elicits so violent a reaction that a boggy nodulopustule, rather like a carbuncle, develops. This is called a kerion, and it usually occurs in a hairy area (e.g. scalp or beard).

Leprosy

During the later stages of lepromatous leprosy, dull, red-brown nodules or plaques may be seen in a symmetrical distribution on the limbs, face and ears. There may be hypopigmented anaesthetic macules elsewhere on the body and impaired eyebrow growth, and the diagnosis is made by finding acid-fast organisms in tissue smears.

Treponema

Late syphilis can produce nodular skin lesions. For example, nodulo-ulcerative tertiary syphilis is characterized by groups of crusted, copper-coloured nodules that spread peripherally and heal centrally in bizarrely shaped patterns. Sometimes, these ulcerate and are seen on the face, trunk or limbs. The solitary syphilitic gumma begins as a dermal or subcutaneous bluish-red nodule. Later, this sloughs to produce a punched-out ulcer with a rubbery necrotic base, and...
NYSTAGMUS

it may cause gross tissue destruction. In secondary yaws, exuberant moist red nodules, rich in Treponema pertenue, may occur widely – particularly in the groins and at the angles of the mouth (their resemblance to raspberries gives rise to the alternative name, ‘framboesia’). Juxta-articular nodules can occur in the tertiary stage of yaws.

Other infections
Leishmaniasis (Baghdad boil) is common in young adults and children in the Middle East, but it can also be seen in European holidaymakers on their return from the south shore of the Mediterranean and the Middle East. Pruritic papules slowly develop into ulcerating nodules on exposed sites following the bite of an infected Phlebotomus fly. Giemsa staining of material readily reveals the intracellular parasite. Onchocerciasis (see p. 528) gives rise to pruritus and indolent non-tender nodules varying in size from 0.5 to 5 cm on the head, shoulders and trunk. These nodules contain the adult worm. In loiasis, or Calabar swelling, transient hot, skin-coloured nodules occur on the face and extremities. Tick bites can be the cause of remarkably persistent dermal nodules.

MISCELLANEOUS
Heberden’s nodes, which are small bony swellings on the terminal interphalangeal joints, are more common in women and are a sign of osteoarthritis. Chondrodermatitis nodularis chronica helicis is the descriptive title given to a not uncommon exquisitely painful nodule that occurs in the upper third of the helix of the ear. The lesions are probably due to pressure, as with a corn, causing underlying perichondritis with fibrinoid degeneration of the cartilage.

NYSTAGMUS

David Werring & Mark Kinirons

Nystagmus refers to an involuntary rhythmic to and fro oscillatory movement of the eyes due to a disorder of the mechanisms responsible for maintaining steady gaze. Nystagmus may result from a dysfunction of the visual system, vestibular apparatus, vestibular nerves, brainstem or cerebellum. There are three mechanisms for maintaining steady gaze: when the eyes are in the primary position, the fixation system is responsible; when the eyes are turned to an eccentric gaze position (e.g. looking to the left), the neural integrator or eccentric gaze-holding mechanism maintains gaze; and when the head is being rotated or looking at a moving object, the vestibulo-ocular system is involved. A problem with any of these systems will cause the eyes to repeatedly drift away from the desired position, leading each time to a corrective saccade that leads to nystagmus.

FIXATION SYSTEM
Fixation involves the ability to detect an eye drift and then initiate an appropriate corrective movement of the eye to keep the desired object on the macula. This requires intact visual input via the optic pathways as well as a complex network involving the parieto-occipital eye fields and the brainstem.

THE NEURAL INTEGRATOR
The neural integrator is responsible for sustaining the eyes at a desired eccentric position in the orbit against the mechanical opposing forces of the orbital tissues. Maintaining eccentric gaze requires a tonic contraction of the extraocular muscles. This ability is impaired by central disorders of the cerebellum or brainstem, or drug intoxication, for example with anticonvulsants. The important brainstem nuclei that constitute this system include the nucleus prepositus hypoglossi, the medial vestibular nucleus for horizontal gaze, the interstitial nucleus of Cajal for vertical gaze, and the vestibulocerebellum. The cerebellar component of the gaze-holding system tends to move the eyes towards the active side of the cerebellum, so that, if there is damage to one side, the eyes tend to drift away from the lesioned side, explaining why the fast phase of nystagmus due to cerebellar damage is towards the side of the lesion and tends to be maximal in this direction.

THE VESTIBULO-OCULAR SYSTEM
The vestibulo-ocular reflex produces eye movements to compensate for head movements, allowing clear vision when the person is moving. Each peripheral vestibular apparatus horizontal semi-circular canal has a tonic input to the brainstem tendency to push the eyes over towards the opposite side. When one side is damaged, the contralateral vestibular apparatus is unopposed and drives the eyes towards the lesioned side, creating the abnormal drift (slow phase) towards the lesion, with corrective jerks (fast phase) away from the lesion. Eye movements away from the lesion will be weakened because of the tonic drift towards the lesion, so when the eyes are moved in the direction of the fast phase, the nystagmus becomes more prominent (Alexander’s law).

DESCRIBING NYSTAGMUS
There are two main types of nystagmus – jerk and pendular – both of which can be horizontal or vertical. Jerk nystagmus consists of a slow phase drifting away from the fixation target, followed by a fast corrective
Nystagmus in childhood

**Congenital nystagmus**

Typically, this condition arises in the first 3 months of life. There may be a clear pattern of inheritance. The nystagmus is horizontal, in all gaze positions (including the primary position), of the jerk or pendular type, and worsened by attempted fixation. About one in six patients will also have eye deviation (strabismus). It may be associated with visual impairment, sometimes with ocular albinism. Brainstem disease can also cause this presentation, so full investigation may be necessary. Latent nystagmus is manifest only when one eye is covered.

**Acquired childhood nystagmus**

Any condition reducing visual acuity in early childhood may cause nystagmus. If acuity is lost before 2 years of age, nystagmus is invariably; if lost between 2 and 6 years, it may develop; if the visual loss is after 6 years old, nystagmus will not usually occur as a result of the visual loss. There are many possible causes, including retinal disorders (e.g. rod or cone dystrophy) or visual pathway damage (e.g. optic chiasm tumour, hereditary optic nerve atrophy or ocular albinism). Extensive investigation may be required, including retinal electrophysiology. Optic nerve glioma is a rare cause of highly asymmetrical or uniocular nystagmus. *Spasmus nutans* refers to a syndrome of nystagmus, head nodding and abnormal head posture. It usually develops between 4 and 12 months of age, and spontaneously resolves by 3 years.

**Acquired horizontal jerk nystagmus**

This occurs when attempting to maintain an eccentric gaze position, and it is the most common type of nystagmus seen clinically. The important distinction is between a peripheral cause due to vestibular dysfunction, and a central cause due to disease of the brainstem or its connections (the neural integrator). This is important because, if a central cause is suspected, urgent neuroimaging must be performed to determine the cause. There are several ways to distinguish between these possibilities:

- The nystagmus of peripheral vestibular dysfunction is (to some extent) suppressed by visual fixation.
- In vestibular dysfunction, the fast phase is always directed away from the damaged ear, and this remains true in both directions of horizontal gaze (i.e., it is unidirectional); the nystagmus is maximal looking away from the lesion.
- In peripheral vestibular damage, the oculocephalic reflex (head impulse test) will be impaired.

**CLINICAL TYPES OF NYSTAGMUS**

To examine for nystagmus, the eyes should be examined at rest, fixating on a target straight ahead (the primary position). The patient should then be asked to follow an object, for example the examiner’s index finger, to the extremes of lateral, upward and downward gaze. Note should be made of whether the pursuit movements are smooth or jerky. Any nystagmus should be carefully noted: whether it is jerk or pendular, its direction, and in which gaze positions it occurs. If there are eye movements at rest, do not forget to look in the mouth to look for evidence of oculopalatal tremor, a rare delayed complication of a brainstem lesion (usually infarction or haemorrhage) interrupting the dentato-olivary pathways (*Mollaret’s triangle*). The patient should then be asked to look in each direction of gaze to assess saccades. Note should be made of any deviation of eye position at rest (strabismus). The vestibulo-ocular reflex or doll’s eye reflex should be tested using the rapid head impulse test: the patient fixates ahead, and the head is rotated rapidly to one side. Intact peripheral vestibular function should mean that the eyes continue to point forwards due to brainstem input resulting from the sudden movement of fluid in the lateral semi-circular canals. An impaired head impulse test provides evidence of peripheral vestibular dysfunction, so if the test is normal, a central cause for symptoms must be considered. A full neurological examination should be performed, including visual acuities, fundoscopy (during which very small amplitude oscillations can be seen), cerebellar and pyramidal functions. Nystagmus may occur in normal subjects at extremes of gaze or with repeated testing, but it usually dampens after a few seconds. Conventionally, three beats is deemed within physiological limits. Signs of congenital disorder, for example ocular albinism, should be sought.

**EXAMINATION OF THE PATIENT WITH NYSTAGMUS**

Phase (a saccade) back towards the target. The direction of the nystagmus is conventionally defined by the fast phase. In pendular nystagmus, both phases of the nystagmus are of the same velocity. Pendular nystagmus is always due to central (brainstem or cerebellar) dysfunction. Nystagmus can be horizontal, vertical or torsional (where the eyes rotate).

Opsoclonus–myoclonus is a dramatic form of abnormal eye movements that may be confused with nystagmus. It is characterized by spontaneous, chaotic, multidirectional saccades. It is often associated with myoclonic limb jerking and is discussed in MYOCLONUS (p. 436).
In peripheral vestibular disturbance, there is often a torsional component.

In peripheral vestibular nystagmus, the patient is usually unable to walk due to severe vertigo and disequilibrium, and will often have nausea, vomiting, sweating or diarrhoea (vegetative symptoms).

In nystagmus due to central disease (causing weakness of the gaze-holding system), the eyes tend to drift back to the central position, causing the fast phase to change depending on the gaze direction, generally being in the direction of the gaze (i.e. it is bidirectional).

Central nystagmus is maximal towards the lesion.

Peripheral acquired horizontal jerk nystagmus
The common peripheral causes of acquired horizontal jerk nystagmus are vestibular neuritis (labyrinthitis) and benign positional paroxysmal vertigo (BPPV). Most patients with vestibular neuritis (perhaps better termed ‘acute peripheral vestibulopathy’, a label that does not imply a definite aetiology) will be prostrated with vomiting, nausea and dizziness without hearing loss, facial weakness or other brainstem dysfunction. The symptoms resolve within a week or so, but may recur. Vestibular sedatives such as cinnarizine may be helpful in reducing symptom severity. BPPV is a symptom complex implying benign end-organ disease. The patient reports brief (less than 40 seconds) vertigo on sudden movements of the head, classically on turning over in bed or lying down. Symptoms ameliorate to some degree when the head is still. There may be a history of viral illness (e.g. upper respiratory tract infection) associated with severe vertigo, or of head trauma. Symptoms can be reproduced by lying the patient supine with their head hanging back over the end of the examination couch (the Hallpike manoeuvre). Treatment with Cawthorne-Cooksey exercises (vestibular rehabilitation) may be helpful.

Drugs may damage the vestibular nerve or end-organ; the main cause is the aminoglycosides including streptomycin and gentamicin.

Ménière’s disease is characterized by recurrent episodes of severe vertigo, vomiting and tinnitus on a background of fluctuating but progressive hearing loss.

Central causes of acquired horizontal jerk nystagmus
Central causes of acquired horizontal jerk nystagmus typically cause less severe vertigo and vegetative symptoms than peripheral causes. Any cause of brainstem or cerebellar disease may be responsible. The acute onset of symptoms may be due to trauma or brainstem or cerebellar infarction; subacute onset may be due to demyelination (multiple sclerosis); slower onset may be due to tumour (either intrinsic, e.g. brainstem glioma, or extrinsic, e.g. cerebellopontine angle tumour). Patients with a cerebello-pontine angle lesion may have nystagmus with the characteristics of both peripheral and central nystagmus. Drugs, for example anticonvulsants (phenytoin) and alcohol, are an important cause of central horizontal jerk nystagmus.

In normal subjects, caloric stimulation induces vestibular nystagmus. Cold water produces horizontal jerk nystagmus to the opposite side, warm water to the same side (mnemonic COWS – Cold: Other; Warm: Same).

Special types of nystagmus
Downbeat nystagmus
This is jerk nystagmus where the fast phase is downwards, usually present in the primary position. The best way to elicit this clinically is in eccentric downward and lateral gaze positions, which typically increase the oscillations. It is an important clinical sign as it has some localizing value. Downbeat nystagmus classically occurs in structural craniocervical junction disease, for example Arnold–Chiari malformation or Paget’s disease. It may also result from cerebellar degenerations (spinocerebellar ataxias or paraneoplastic cerebellar degeneration), brainstem disease such as multiple sclerosis, glioma or infarction, drugs (lithium and alcohol), brainstem encephalitis, magnesium depletion or Wernicke’s encephalopathy.

Upbeat nystagmus
Jerk nystagmus with a fast phase upwards, usually present in the primary position, suggests structural disease of the cerebellar vermis or brainstem (e.g. a posterior fossa tumour).

See-saw nystagmus
This is a striking disorder in which one eye elevates and intorts while the other eye drops and extorts. It occurs with abnormalities at the junction of the mesencephalon and diencephalons, for example a tumour.

Convergence-retraction nystagmus
This is characterized by rhythmical convergence or retraction movements of the eyes when the patient tries to look up. It is a component of Parinaud’s syndrome together with light–near dissociation, lid retraction, vertical gaze paresis and impaired accommodation. This syndrome is due to pathology in the dorsal midbrain at the level of the superior colliculus. A pineal gland tumour (pinealoma) is a classic cause.
**Periodic alternating nystagmus**

This is spontaneous horizontal jerk nystagmus that reverses every 90–120 seconds with a 1–10-second pause between reversals. Its localizing significance is similar to downbeat nystagmus. It may respond to baclofen.

**Voluntary nystagmus**

A few people can perform very high frequency horizontal saccades that resemble pendular nystagmus. The eyelids often flutter at the same time, and the movements usually fatigue after more than a few minutes.
disordered.

others occurs, the personality may be considered individual difficult to live with. Once harm to self or work, but sometimes are so marked as to make the personality disorder. Here they are referring to traits perfectionism and rigidity. These can all be adaptive personality traits and obsessional (or anankastic) finally, one other usage of the term ‘obsessional’ must be mentioned. Psychiatrists talk about obsessional conceptualized as a variant of OCD. The term ‘dependent oedema’ has, therefore, no specific diagnostic significance, and its only value is as a reminder to beginners to look for oedema in those parts of the body which are dependent at that time. Thus, a decrease in ankle oedema after a night’s rest is not necessarily a sign of improvement; in such circumstances, the oedema may well have moved and be found in the back and sides of the calves and thighs. This behaviour of the oedema fluid accounts for an occasional curious finding such as rings of oedema around the olecranon processes; this can be due to the patient spending much of his time in bed leaning forward with his elbows on a bed table in front of him. The causes of oedema, nearly always dependent, are discussed under OEDEMA, GENERALIZED (below).

Generalized oedema is due to an increase in the volume of extracellular fluid. This is brought about by excessive renal tubular reabsorption of sodium and water; the mechanism of this reabsorption is complex, and the renin–angiotensin–aldosterone system is only one of the factors involved. The accumulation of fluid in the extravascular space that causes oedema is determined by the relationship between the hydrostatic and oncotic pressures in the capillaries and the interstitial tissue. Thus, a rise in capillary hydrostatic pressure due, for example, to venous obstruction and a fall in capillary oncotic pressure, as a result of hypoalbuminaemia, increases the net movement of fluid from the capillaries to the tissues. When sufficient fluid has been transferred in this way – at least 5 litres in adults – clinically detectable oedema results. Another factor concerned in the transfer of fluid across capillary walls is their permeability; in practice, an increase in this is more important in the production of localized rather than generalized oedema. Impairment of the lymphatic drainage of tissues is also a cause of localized oedema (see OEDEMA OF LEG, p. 472).
In any patient with generalized oedema, fluid may also accumulate in serous cavities in the form of ascites and pleural or pericardial effusion.

Gravity determines the fact that, in any situation in which sodium and water retention occurs, the oedema fluid tends to accumulate in the dependent parts of the body (see OEDEMA, DEPENDENT, p. 470). This tendency is so marked that, even in normal subjects, a little ankle oedema is common following prolonged periods of immobility in the seated position. It is particularly common during long journeys by air and, sometimes, by train or coach; it is probable that this is mainly due to a reduction in lymphatic drainage, which is critically dependent on muscular activity. Also, in about 90 per cent of pregnant women, slight oedema is present at term. The pathological situations in which generalized oedema most commonly occurs are heart failure and renal, hepatic and, less often, gastrointestinal disease. These, and other less common causes of oedema, are discussed below.

HEART FAILURE

The oedema of heart failure is typically dependent, affecting particularly the ankles and, in recumbent patients, the sacral region. Despite this clear evidence of a hydrostatic component in determining the site of the oedema, the rise in central venous pressure in heart failure is a minor factor in the production of the oedema, compared with the reduction in renal blood flow and glomerular filtration rate, and the increase in tubular reabsorption of sodium and water. These mechanisms operate whatever the cause of the heart failure but, in cor pulmonale, an additional factor may be a movement of fluid from the cells to the interstitial tissue; this is believed to occur in order to provide more buffers for the associated respiratory acidosis. The oedema is usually symmetrical, but it is sometimes more marked in the left leg than the right; this is thought to be due to pressure on the left common iliac vein by the right common iliac artery as it crosses it. It is heart failure that is the mechanism of the oedema of so-called ‘wet beri-beri’, due to a dietary deficiency of thiamine, and also of the oedema seen commonly in the trunk and face as in the legs; the external genitalia are commonly very swollen. Spontaneous disappearance of the oedema is not necessarily a good sign as, with advancing renal failure, the fall in glomerular filtration rate may markedly reduce the amount of protein lost.

There are numerous conditions that can cause the nephrotic syndrome. In children, by far the most common cause is minimal-change nephropathy, in which, as the name implies, the glomeruli are nearly normal on light microscopy but show characteristic changes on electron microscopy; clinically, the most typical feature is the complete remission produced by steroid therapy. In adults, membranous and focal and segmental glomerulosclerosis are more common than the minimal-change lesion. Other common causes include systemic lupus erythematosus, amyloidosis and diabetic nephropathy; oedema is quite common in individuals with diabetes even in the absence of heavy proteinuria, perhaps due to microvascular disease. There is also a recognized association of the nephrotic syndrome with malaria due to Plasmodium malariae, and with malignant disease, especially adeno-carcinoma and lymphoma. HIV-associated nephropathy predominantly affects Afro-Caribbeans and causes severe nephrotic syndrome and rapidly deteriorating renal function. The high venous pressure of constrictive pericarditis has also occasionally been known to cause nephrotic syndrome, which has in addition been seen in cyanotic congenital heart disease. Renal vein thrombosis per se, however, is no longer thought to be a cause but rather a complication of the nephrotic syndrome. A number of drugs and other substances are known to cause a membranous glomerular lesion and proteinuria heavy enough to cause oedema; these include non-steroidal anti-inflammatory drugs, gold, penicillamine and captopril. A specific allergy is probably responsible for the nephrotic syndrome associated with certain foods, pollens, penicillin, bee stings and poison ivy.

RENAL DISEASE

In nephrotic syndrome hypoalbuminaemia, due to heavy proteinuria, is severe enough to lower the intracapillary oncotic pressure to a level at which oedema occurs. In this condition, the urinary protein loss is usually more than 3 g per 24 hours, and the serum albumin below 30 g/l. It is probable that the hypovolaemia resulting from a massive loss of fluid from the capillaries stimulates the renin–angiotensin–aldosterone system, and this leads to renal retention of sodium and water. The oedema is usually dependent but, in children, it may be as prominent in the trunk and face as in the legs; the external genitalia are commonly very swollen. Spontaneous disappearance of the oedema is not necessarily a good sign as, with advancing renal failure, the fall in glomerular filtration rate may markedly reduce the amount of protein lost.

LIVER DISEASE

Fluid retention is common in hepatic failure, in which it is due to impaired protein synthesis and consequent hypoalbuminaemia. The changes in renal function are similar to those in nephrotic syndrome. Both oedema and ascites can occur, often together, but one can be
be little difficulty in distinguishing them from the

In most of these cases, both legs will be involved, but occasionally – and perhaps due to the patient’s position in bed – the oedema may be somewhat asymmetrical. In most of the conditions discussed in this section, the oedema is strictly unilateral, and there will, therefore, be little difficulty in distinguishing them from the

The distribution of the oedema is curious, affecting particularly the face, hands, breasts, thighs, buttocks and abdominal wall, and hardly ever the ankles. The aetiology is unknown, but it has been suggested that abnormal autonomic reflexes may be responsible.

Oedema is one of the cardinal signs of inflammation. Local lesions, such as boils and carbuncles, are easily identified, but more widespread inflammatory lesions may initially cause diagnostic confusion. The bright red areas with palpable raised margins of erysipelas are characteristic, but oedema due to obstruction of cutaneous lymphatics may persist after the acute inflammation has subsided. Cellulitis causes more diffuse oedema, as does acute osteomyelitis; lymphangitis and lymphadenitis are more common in the former while, in the latter, the constitutional disturbance is greater. Chronic osteomyelitis can cause puzzling oedema, but imaging will settle the diagnostic issue. Acute arthritis of any type is associated with local oedema. In most cases, the swelling of the joint itself will make the diagnosis clear, but acute gout can cause a swelling that is widespread enough to simulate cellulitis. Acute rheumatoid arthritis is less likely to cause confusion, but it is worth remembering that, in this condition, more generalized oedema can occur; this is probably due to a combination of hypoalbuminaemia, stasis in an immobile patient and, perhaps, increased capillary permeability. The painful swelling of the calf with ankle oedema caused by a ruptured Baker’s cyst is easily confused with deep venous thrombosis; a history of prior swelling of the joint, decreasing with the onset of the calf pain, is an important diagnostic feature.

In the great majority of patients with generalized oedema, the legs, particularly the ankles, are likely to be affected (see OEDEMA, GENERALIZED, p. 470). In most of these cases, both legs will be involved, but occasionally – and perhaps due to the patient’s position in bed – the oedema may be somewhat asymmetrical. In most of the conditions discussed in this section, the oedema is strictly unilateral, and there will, therefore, be little difficulty in distinguishing them from the

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VENOUS OBSTRUCTION

Deep venous thrombosis in the calf muscles is an important cause of oedema of the ankle; the swelling will extend further up the leg if thrombus is also present in the femoral and iliac veins. Venous thrombosis is a common complication of major surgery and of prolonged recumbency for any reason; it can also often occur without any obvious cause. Long coach journeys by the elderly and long-haul flights and, in women, the use of oestrogen-containing oral contraceptive agents increase the risk of deep venous thrombosis. In addition to the ankle oedema, the calf is typically swollen and tender on pressure from behind and from side to side. Homan’s sign (pain in the calf on dorsiflexion at the ankle) may be positive, but is of no diagnostic value as it can be positive with any painful lesion of the calf. Very often, none of these signs is present; the first evidence of deep venous thrombosis may be a fatal pulmonary embolism.

Thrombus may spread upwards beyond the iliac veins to involve the inferior vena cava and, in that case, the other leg will become oedematous. With such a sequence of events, the diagnosis of inferior vena caval occlusion is clear. Primary occlusion of this vessel, however, will present with bilateral ankle and leg oedema, in which case the differential diagnosis includes all the causes of generalized oedema. Diagnostic assistance may be provided by investigations such as Doppler flow studies, but the definitive diagnosis of venous occlusion can be made only by venography or computed tomography scanning. In any patient in whom venous thrombosis has occurred without obvious cause, a thorough general examination – including rectal and vaginal examination – is essential to detect pelvic or abdominal tumours causing pressure on the veins.

Varicose veins, with or without previous thrombosis, commonly cause ankle oedema, usually quite mild. Although in the great majority of cases it is the veins themselves which are the seat of the trouble, it is wise, at the first presentation, to examine the patient with the possibility of external pressure on the large veins in mind, as in patients with acute venous occlusion.

LYMPHATIC OBSTRUCTION

The oedema fluid in lymphatic obstruction contains much more protein than the fluid in other conditions causing oedema. Consequently, although initially the swelling ‘pits’ on pressure and disappears overnight, it later becomes firm and non-pitting (‘brawny’) and is present all the time. Later, the skin becomes grossly thickened and, sometimes, ulcerated. The common causes of lymphatic obstruction in Western countries include neoplastic infiltration of the lymphatics and lymph nodes, and scarring from trauma, surgical or otherwise, and from irradiation; recurrent streptococcal infection can cause similar damage. In tropical countries, filariasis is a common cause of lymphatic obstruction and chronic oedema of the legs known as ‘elephantiasis’. The most common form of filariasis is that due to *Wuchereria bancrofti*; this is widespread in the tropics and causes elephantiasis of the whole leg and also of the genitalia, especially in males. Another filaria, *Brugia malayi*, is found in the Far East, especially South-East Asia; the inguinal lymph nodes are affected, as in Bancroftian filariasis, but curiously the oedema and elephantiasis are confined to the lower parts of the legs. There is also a non-filarial elephantiasis seen in parts of Africa and Central and South America; this is due to chronic lymphangitis apparently caused by microscopic particles of silica absorbed through the skin of the feet; it usually affects the feet and lower parts of the legs. Lymphatic hypoplasia is a rare cause of oedema that occurs in two forms: a congenital variety presenting with oedema in early infancy; and a familial form, sometimes known as Milroy’s disease. In the latter, there is an abrupt demarcation between the swollen and the normal tissue at the level of a joint – ankle, knee or hip (Fig. O.1). There is sometimes a history of attacks of
OLIGURIA

James Pattison

The volume of urine that constitutes clinically significant oliguria varies with the pre-existing state of the kidneys. With a urine volume of less than 400 ml per 24 hours, even a normal kidney is unable to concentrate the glomerular filtrate sufficiently to prevent a rise in plasma urea and creatinine. Much larger volumes may be needed to maintain homeostasis if renal function has been impaired for any reason. Oliguria is present in most, but by no means all, cases of acute kidney injury (previously known as acute renal failure), of which the causes, summarized in Box 0.1, are conventionally classified as prerenal, renal and postrenal. Although these categories are not entirely mutually exclusive, it is important to identify, in any individual patient, the dominant factor causing acute kidney injury, because the management is very different for these three groups. It is also important to decide whether the condition from which the patient is suffering is acute kidney injury with previously normal kidneys, or an acute exacerbation of chronic renal disease. The latter is more likely if the patient is very anaemic, shows evidence of long-standing hypertension or has biochemical or radiographic evidence of osteodystrophy, or, most significantly, if the kidneys can be shown to be shrunken on ultrasound scanning.

RENNAL CIRCULATORY INSUFFICIENCY (PRERENAL URAEMIA)

This situation arises whenever the renal blood flow falls steeply. It is most commonly due to a fall in cardiac output secondary to a reduction in circulating blood volume. The implication of a diagnosis of prerenal uraemia is that normal renal function will be restored as soon as the circulatory abnormality has been corrected. The cause of the circulatory failure is usually obvious. External loss of fluid is a common cause; severe diarrhoea and vomiting, haemorrhage, burns and previous polyuria in diabetes mellitus or Addison’s disease are all well-known causes of, usually transient, oliguria. Other mechanisms operate in so-called ‘cardiogenic’ shock following myocardial infarction, acute pancreatitis and septicaemia, often due to Gram-negative organisms. All these conditions can also cause acute tubular necrosis, a poorly defined pathological entity, but a diagnosis that is made when renal function does not improve rapidly following restoration of normal renal perfusion. It is clearly important not to delay the recognition that this change for the worse has taken place, and a useful indication may be provided by estimation of the urinary sodium concentration. With oliguria due to renal circulatory insufficiency, this is typically very low, around 20 mmol/l, whereas in acute tubular necrosis with oliguria it will be about three times that level.

RENNAL CAUSES OF OLIGURIA

Acute tubular necrosis

As has been said, this is an imprecise term and it will be used here to designate only those cases of acute, usually oliguric, kidney injury due to those circulatory disorders described above under prerenal uraemia. Other specific disorders causing a similar clinical syndrome will be discussed separately. In practice, the diagnosis will usually be made in patients under observation and treatment for the causative condition. The symptoms are those of uraemia, i.e. anorexia, nausea and vomiting, perhaps muscle cramps and a ‘flapping’ tremor. Bleeding into the skin and gastrointestinal tract and fits can occur. Hypertension is unusual and suggests a different cause for the renal failure. The plasma creatinine will rise progressively unless effective treatment is started, and dangerous hyperkalaemia, especially when there has been much tissue destruction, is common. Recovery can usually be expected for up to 6 weeks; if this does not occur, renal biopsy may reveal that the damage is severe, perhaps in the form of acute cortical necrosis. In such a case, recovery is very slow and may be incomplete.

Box 0.1 Causes of oliguria

- Renal circulatory insufficiency
- Primary renal disease
  - Acute tubular necrosis
  - Acute cortical necrosis
  - Drugs and poisons
  - Acute interstitial nephritis
  - Acute glomerulonephritis
  - Vascular lesions (e.g. accelerated hypertension, haemolytic uraemic syndrome)
  - Myoglobinuria secondary to rhabdomyolysis
  - Haemoglobinuria secondary to intravascular haemolysis
- Obstructive
  - Calculi
  - Pelvic tumours
  - Retroperitoneal fibrosis
Acute cortical necrosis
This condition may follow insults similar to those which cause acute tubular necrosis. It is a recognized feature of severe obstetric emergencies such as antepartum haemorrhage, eclampsia and septic abortion. Irreversible renal damage occurs, but the changes may be patchy and, therefore, compatible with some recovery. Radiography may show shrunken kidneys with cortical calcification as early as 2 months after the onset.

Acute renal failure due to drugs and poisons
Many drugs are known to damage the kidneys and cause acute kidney injury. Some do so by causing acute interstitial nephritis (see below). Others, such as the aminoglycoside antibiotics, amphotericin, colistin, polymyxin B and radiographic contrast media, do so by other mechanisms. Heavy metals, organic solvents, particularly carbon tetrachloride, paraquat, snake bites and mushroom poisoning can also cause acute renal failure.

Acute interstitial nephritis
This is a common cause of acute oliguric kidney injury and is most often due to drugs. Those most commonly implicated include the non-steroidal anti-inflammatory drugs (NSAIDs), the penicillin and cephalosporin groups of antibiotics, and diuretics, but many other drugs are known occasionally to cause this syndrome. The same type of renal damage is occasionally caused by bacterial and viral infections.

Acute glomerulonephritis
Some reduction in urine volume is common in most cases of acute nephritis, and severe oliguric acute kidney injury is a common feature, particularly in adults, of the more rapidly progressive forms of glomerulonephritis, which are discussed under HAEMATURIA (p. 241).

Vascular lesions
Renal infarction, if extensive, may cause acute renal failure; renal vein thrombosis is much less likely to do so except in infants. The fibrinous necrosis of accelerated (malignant) hypertension, as well as the similar vascular damage seen occasionally in systemic sclerosis, together with intravascular coagulation occurring in the haemolytic–uraemic syndrome of children, thrombotic thrombocytopenic purpura and the rare idiopathic postpartum renal failure, are rare causes of oliguria and renal failure.

Other causes of acute oliguric kidney injury
Acute kidney injury may complicate hepatic failure from any cause; this hepatorenal syndrome is thought to be related to alterations in blood flow induced by the hepatic failure rather than intrinsic renal disease per se. Muscle damage is an important contributory cause of renal failure following trauma, and rhabdomyolysis in the absence of trauma can also cause renal damage. This can be due to acute myositis, either idiopathic or in association with viral infections, prolonged convulsions or marathon-running, malignant hyperpyrexia following general anaesthesia, carbon monoxide poisoning and a number of other conditions. It is probably the myoglobinuria which is responsible for the renal damage, but the mechanism by which this occurs is unclear. Similar damage can occur as a result of intravascular haemolysis, as in malignant malaria or following a mismatched blood transfusion. In myelomatosis, acute renal failure may occur, probably due to hypercalcaemia as well as the specific renal lesion of that condition. Finally, obstruction of the renal tubules by crystals of urate, as part of tumour lysis syndrome during the treatment of lymphoma and similar conditions, is a rare but important and preventable cause of acute oliguric renal failure.

POSTRENAL (OBSTRUCTIVE) CAUSES OF Oliguria
Unlike primary renal disease, obstructive lesions below the renal papillae more commonly cause total anuria than oliguria, and this is sometimes a helpful differential diagnostic feature. Obstructive anuria can occur only if the outflow from both kidneys or from the only functioning kidney is obstructed. In a patient with acute renal failure and anuria, or severe oliguria, it is clearly important to distinguish between an obstructive cause, which can usually be quickly dealt with surgically, and a primary renal cause, for which other treatment is required. The situation is not always clear-cut. For example, chronic stone disease may cause severe renal damage and chronic renal failure without much in the way of symptoms, as well as acute anuria from obstruction; or chronic ureteric obstruction by spread from a uterine carcinoma might be complicated by pyonephrosis and Gram-negative septicemia causing acute tubular necrosis.

Stone disease is the most common cause of obstructive anuria. It is rare for this to occur as a result of simultaneous obstruction of both ureters by calculi; it is more common to find that one kidney has been severely damaged by chronic disease, and that there is a calculus in the other ureter. If the patient has total anuria, it is necessary to confirm the diagnosis and localize the obstruction. Radiographic examination by ultrasound of the kidneys and ureters is essential, with CT KUB (kidneys, ureters and bladder) to demonstrate a small or poorly radio-opaque calculus.
Other causes of anuria from ureteric obstruction include bladder carcinoma; in such cases, it is likely that the diagnosis will already have been made, and chronic partial bilateral ureteric obstruction will have caused hydroureter and hydronephrosis with severe renal damage. A similar situation can arise in carcinoma of the uterine cervix, but anuria may occasionally be the presenting feature. Other pelvic or abdominal tumours can also cause bilateral ureteric obstruction, and ligation of both ureters is an occasional complication of extensive pelvic surgery for malignant disease. Retroperitoneal fibrosis is a rare cause of bilateral ureteric obstruction; in most cases this is now thought to be due to IgG4 disease, but it can also be caused by retroperitoneal lymphoma, and peri-aortitis around an atherosclerotic aorta. Sulphonamide crystalluria with blockage of both ureters is now very rare, although the need for a high fluid intake in patients on these drugs remains.
Considerable difficulty may be experienced in viewing a highly myopic fundus even through a dilated pupil, although the patient’s own spectacles in situ reduce magnification and provide a better view. When the condition is marked and associated with distortion of the disc itself, it is referred to as disc coloboma; astigmatism or hypermetropia is then often marked and the vision reduced, even with correction of the refractive error.

PIGMENTATION
The disc margin is variably pigmented, from a small crescent to a complete ring. The pigment has no pathological significance. The degree of pigmentation of the fundus varies considerably, generally corresponding to the complexion of the patient. When there is little or no retinal epithelial pigment, the choroidal vessels are plainly seen (albinoid fundus); when retinal pigmentation is intense, the appearance of tigroid fundus is presented. The retina may also show small congenital oval or round greyish-black pigment spots.

MYELINATED NERVE FIBRES
This normal variant can be recognized by the brilliant white colour of the myelinated nerve fibres, the feathered, striated appearance, and the embedding of the retinal vessels among the nerve fibres (Fig. O.5).

DRUSEN (COLLOID BODIES)
These are seen as multiple oval or round, yellowish spots, especially around the optic disc. They are common in later life, and usually do not affect vision. They may be confused with papilloedema. Calcified drusen can be detected on computed tomography or ultrasound scanning.

PSEUDOPAPILLOEDEMA
In marked hypermetropia the disc may be small, crowded and elevated, with no physiological cup and an ill-defined margin; the vessels may be tortuous (although not dilated) and, unless the error of refraction is observed, the condition may be mistaken for papilloedema. In true disc swelling, the vascular changes are usually prominent, with capillary dilatation, exudates and haemorrhages. If spontaneous venous pulsation of the central retinal vein is seen, raised intracranial pressure (ICP) is virtually excluded – although the absence of pulsation is common in normal individuals and is not a helpful indicator of raised ICP. Flourescein angiography may be helpful in distinguishing pseudopapilloedema from true swelling.

COLOBOMA OF THE CHOROID
This is a congenital defect that may be recognized by its situation, which is usually below the disc, as an oval area of exposed sclera that may extend from the periphery and include the optic disc, with overlying healthy retinal vessels on its surface (Fig. O.6). It may be associated with other congenital abnormalities such as coloboma of the iris or lens.

ABNORMALITIES OF THE OPTIC DISC
Optic disc swelling (papilloedema)
Although the terms ‘optic disc swelling’ and ‘papilloedema’ are often used interchangeably, papilloedema is conventionally applied specifically to disc swelling (usually bilateral) due to raised ICP. The underlying cause of any optic disc swelling is stasis of retrograde axonal transport along the optic nerve. This may be due to a variety of different pathologies. Raised ICP increases pressure in the anterior optic nerve sheath, impairing axonal function at the
lamina cribrosa; in anterior ischaemic optic neuropathy, an impaired blood supply to the nerve head disrupts axonal transport. In optic neuritis, inflammatory cell accumulation around vessels, with interstitial oedema, compresses axons at the optic nerve head.

Swelling of the disc causes loss of the normal physiological cup and blurring of the disc margin. The retinal veins are often dilated, and there may be flame-shaped haemorrhages on the disc surface. The main causes of optic disc swelling are listed in Box O.2.

In the differential diagnosis of optic disc swelling, it is helpful to distinguish between disc swelling due to raised ICP and that due to local disease affecting the optic nerve. A raised ICP usually causes bilateral, symmetrical optic disc swelling, with acuity and colour vision usually preserved (Fig. O.7). Local pathology of the optic nerve usually causes unilateral disc swelling with early loss of acuity and colour vision, for example in anterior optic neuritis or ischaemic optic neuropathy.

Papilloedema due to raised ICP has a characteristic natural history: early (blurred margins and loss of venous pulsation); acute (secondary haemorrhages, hard exudates and cotton-wool spots); chronic (swelling with mild venous congestion); vintage (early disc pallor); and finally atrophic (established optic atrophy). When the atrophy follows papilloedema (Fig. O.8), there may be glial sheathing of the vessels at the disc (secondary optic atrophy), oedema residues may fill the physiological cup, the colour is then greyish-white, the retinal vessels are thin and tortuous, and the edge of the disc is irregular.

Optic disc cupping

Excavation of the optic disc (cupping), occurs in chronic glaucoma (see Fig. O.3). There is enlargement of the cup associated with thinning of the normally pink neuroretinal rim with associated peripheral

**Box O.2  Differential diagnosis of optic disc swelling**

- **Raised intracranial pressure** (usually bilateral)
  - Tumour
  - Abscess
  - Haematoma
  - Cerebral venous sinus thrombosis
  - Meningitis (granulomatous, carcinomatous, haemorrhagic)
  - Subarachnoid haemorrhage
  - Cerebrospinal fluid-secreting tumour (choroid plexus tumour) – rare
  - Malignant hypertension

- **Local pathology** (usually unilateral – unless the disease affects both optic nerves at the same time)
  - Anterior optic neuritis
  - Anterior ischaemic optic neuropathy
  - Optic nerve compression by tumour (e.g. glioma, lymphoma)
  - Optic nerve infiltration (e.g. granulomatous disease – tuberculosis, sarcoid)
  - Central retinal vein occlusion

**Figure O.6** Coloboma of the choroid.

**Figure O.7** (a) Acute papilloedema; (b) chronic papilloedema.
visual field loss. The cupping is particularly evident in the vertical meridian. The cupping of the optic disc in cases of glaucoma may be distinguished from the physiological cup by the fact that it extends beyond the physiological norm – that is, more than 60 per cent of the size of the disc in the vertical meridian. The retinal vessels bend sharply over the edge and may disappear from view behind the overhanging margin of the disc, reappearing on the base of the cup. The lamina cribrosa is clearly seen and the disc becomes white and atrophic.

**Optic atrophy**

Pallor of the disc (Fig. O.9) signifies atrophy of the disc (i.e. a reduction in the number of nerve fibres that arise in the retina and converge to form the optic disc). The causes are listed in Box O.3.

**Peripapillary atrophy**

Atrophy of the tissues around the disc occurs with age. In younger patients, it is most frequently seen with myopia (i.e. myopic crescent) (see Fig. O.4). This is usually found on the temporal side of the disc and may vary in size and extent from a thin crescent to a large atrophic area around the whole disc.

**Retinal vascular abnormalities**

**Central retinal vein occlusion**

This condition makes the disc extremely swollen and oedematous, with dramatic blurring of the edge (Fig. O.10). There may be macular oedema. The retinal veins are dilated and tortuous, and the whole fundus may be covered with flame-shaped and blotchy haemorrhages. The oedema of the retina may entirely obscure retinal venous segments. Cotton-wool spots may be seen. The causes and associations are listed in Box O.4.
Distal to the occlusion, there are numerous scattered haemorrhages, dilated tortuous veins and retinal oedema. Central retinal vein occlusion is associated with severe visual loss with little prospect of recovery, while branch retinal vein occlusion has a much more favourable prognosis.

Central retinal artery occlusion

This condition occurs more commonly in the middle-aged or elderly. Visual loss occurs following loss of blood supply to the inner layer of the retina (Fig. O.1). The central retinal artery is the first intraorbital branch of the ophthalmic artery, which enters the optic nerve behind the globe to supply the retina. Short posterior ciliary arteries arise more distally from the ophthalmic artery and supply the choroid.

Cardiac embolism must always be considered, especially in young patients who may have valvular disease, right-to-left shunting from patent foramen ovale, or endocarditis. The emboli are usually cholesterol, but they may be bacterial or talc in some patients (e.g. intravenous drug users). In older patients (over 50 years), carotid atherosclerosis and thromboembolism, and giant-cell arteritis, are the most important causes of retinal artery occlusion. The main causes of central retinal artery occlusion are listed in Box O.5.

The infarcted retina is pale due to secondary oedema. The central macula (fovea) is spared because it has additional supply from the choroidal capillary circulation, and it appears as a bright ‘cherry-red’ spot. The differential diagnosis of a cherry-red spot includes a number of lipid storage diseases (Box O.6), which are characterized by more widespread neurological dysfunction.

No visual recovery is likely following central retinal artery occlusion. The pallor of the inner retina slowly clears, and secondary atrophy of the optic disc later develops. If giant-cell arteritis is suspected (a history of preceding amaurosis, age over 60 years, constitutional symptoms and high erythrocyte sedimentation rate), urgent treatment with high-dose corticosteroids may save vision in the other eye.

Investigation will depend on age and clinical presentation, but it may include carotid or cardiac imaging to seek sources of emboli.

Renal retinopathy

Renal retinopathy is characterized by the presence of flame-shaped haemorrhages in the nerve fibre layer.
of the retina. There are also two types of white patch: in the early stages of the disease, ill-defined ‘cotton-wool spots’ and, in later stages, smaller linear patches of white exudate may be seen radiating from the macula.

**Hypertensive retinopathy**

The arteries are narrow, show a heightened light reflex (silver or copper wiring), and compress the veins at the crossings (arteriovenous nipping), which may show local deflections at these points. In more severe hypertension, there is retinal ischaemia and loss of the integrity of the small vessels; this results in the appearance of retinal exudates (due to oedema) and ‘cotton-wool spots’, which can be confused with diabetic retinopathy (see below). Cotton-wool spots are white, fluffy lesions that result from occlusion of precapillary arterioles supplying the retinal nerve fibre layer, with subsequent swelling of nerve fibres (Fig. O.12). They are also called ‘soft exudates’ or ‘nerve fibre layer infarctions’. Fluorescein angiography shows no capillary perfusion in a cotton-wool spot. Other causes of cotton-wool spots are listed in Box O.7. Disc swelling is also evident in malignant hypertension. Hard exudates (intraretinal lipid exudates) result from the leakage of lipid through damaged capillaries, and these may be visible as well-defined yellow deposits within the retina, sometimes in a circinate pattern. Hyperlipidaemia may correlate with the development of hard exudates.

**Diabetic retinopathy**

This frequent complication of long-standing diabetes mellitus is a common cause of blindness in adults in the Western world. The complex damage to the retinal microcirculation causes a number of characteristic features, including microaneurysms, dot and blot haemorrhages, neovascularization, cotton-wool spots and hard exudates (Fig. O.13). Microaneurysms (focal dilatations) of abnormal retinal capillaries are seen as small red dots. Visual loss may occur due to leakage of lipids and water from abnormal capillaries into the retina. Deep retinal haemorrhages occur when vessel walls or aneurysms rupture (‘dot’ or ‘blot’ haemorrhages). More superficial haemorrhages may be flame-shaped and indistinguishable from those of hypertensive retinopathy. Retinal ischaemia due to widespread capillary occlusion or hypoperfusion results in the production of vasoproliferative substances and in neovascularization. Neovascularization can involve the retina, optic disc or the iris, the latter being an ominous sign of severe proliferative disease that may be complicated by intractable glaucoma. The new retinal vessels bleed easily, causing vitreous haemorrhage (Fig. O.14) and may give rise to fibrous proliferation, which may cause traction retinal detachment. Nerve fibre layer infarcts (cotton-wool spots) and hard exudates are also features of diabetic retinopathy, particularly in patients who are also hypertensive.

**Retinal vasculitis**

There are many systemic associations with retinal vasculitis, which is characterized by perivascular sheathing by inflammatory cells. The important causes are listed in Box O.8.

Figure O.12 Hypertensive retinopathy.

**Box O.7  Differential diagnosis of cotton-wool spots**

- Pre-proliferative diabetic retinopathy
- Hypertension
- Retinal vein or branch retinal artery occlusion
- HIV retinopathy
- Autoimmune disorders (e.g. systemic lupus erythematosus)
- Scleroderma
- Haematological disorders

Figure O.13 Diabetic retinopathy.
Macular diseases

Age-related macular degeneration

This is the most common cause of blindness in Western countries. It is due to age-related changes in the retinal pigment epithelium (the most metabolically active part of the retina), and small choroidal vessels. The fundus changes include atrophy and clumping of the pigment and atrophy and sclerosis of the choroidal blood vessels. Drusen (hyaline deposits) between the retinal pigment epithelium and choroid also appear (Fig. O.15). In some patients, sudden visual loss results in haemorrhage in subretinal neovascularization. Occasionally, laser treatment may halt the progress of subretinal neovascularization, but otherwise no treatment is available for this condition. Patients often benefit, however, from low visual aids.

Box O.8  Causes of retinal vasculitis

<table>
<thead>
<tr>
<th>Category</th>
<th>Conditions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Idiopathic</td>
<td>Giant-cell arteritis</td>
</tr>
<tr>
<td>Infectious</td>
<td>Scleroderma</td>
</tr>
<tr>
<td>Herpes (zoster or simplex)</td>
<td>Other systemic diseases</td>
</tr>
<tr>
<td>Cytomegalovirus</td>
<td>Sarcoïdosis</td>
</tr>
<tr>
<td>HIV</td>
<td>Behçet’s disease</td>
</tr>
<tr>
<td>Syphilis</td>
<td>Multiple sclerosis</td>
</tr>
<tr>
<td>Tuberculosis</td>
<td>Malignancy</td>
</tr>
<tr>
<td>Toxoplasmosis</td>
<td>Crohn’s disease</td>
</tr>
<tr>
<td>Collagen vascular diseases</td>
<td>Churg–Strauss syndrome</td>
</tr>
<tr>
<td>Systemic lupus</td>
<td>Whipple’s disease</td>
</tr>
<tr>
<td>erythematous</td>
<td>Miscellaneous</td>
</tr>
<tr>
<td>Polyarteritis nodosa</td>
<td>Eales’ disease</td>
</tr>
</tbody>
</table>

Macular oedema

Macular oedema may be caused by many disease processes, all of which have in common a breakdown of the blood–retinal barrier with an increase in extracellular fluid within the retina. The retinal capillary endothelium and the retinal pigment epithelium form this barrier, and diseases affecting either may cause oedema with accompanying visual loss. The main causes are retinal vascular disease (e.g. retinal vein occlusion and diabetic retinopathy).

Peripheral retinal abnormalities

Retinal detachment

The retina may become detached from the underlying retinal pigment epithelium by a variety of mechanisms. Most commonly, fluid vitreous tracks through a break or tear in the retina, progressively lifting the retina around the break. The area of detachment slowly spreads until the whole retina is detached (Fig. O.16). Unless the retinal break is sealed surgically, vision will be lost. Retinal breaks are most common in myopic patients and may follow trauma. The second mechanism is when traction forces from the vitreous pull the retina off. This may occur under conditions where there is fibrous proliferation within the eye, for example in advanced diabetic retinopathy or following penetrating injuries of the eye. Finally, the retina may detach because it is pushed off either by a tumour of the underlying choroid or by subretinal exudate, which may complicate inflammatory choroidal disease or vascular anomalies. The detached retina has a grey appearance, and the retinal vessels are tortuous and thin, and appear dark in colour.
Retinitis pigmentosa
This is a group of hereditary diseases affecting the photoreceptors and retinal pigment epithelium. Night blindness and constriction of the visual fields (tunnel vision) are variably progressive, with eventual blindness. The earliest fundus changes occur in the equatorial zone with areas of hyperpigmentation (typically in a bone spicule pattern), which spread to involve the whole fundus. Waxy pallor of the disc and gross attenuation of the retinal arterioles are also features (Fig. O.17).

Choroiditis
Active choroiditis manifests itself as a greyish-white, ill-defined, slightly raised area. Overlying vitreous opacities are commonly associated with such areas of active choroiditis. When healed, the patch appears white because there is atrophy of the choroid and overlying retinal pigment epithelium. Such scars are generally surrounded by clumps of dense pigment (Fig. O.18).

Eales’ disease
This rare disorder manifests itself as recurrent intraocular haemorrhages in young adults and is generally attributed to retinal phlebitis. Sudden obscuration of vision results from haemorrhage into the vitreous. Recurrence is the rule, and proliferative retinopathy may ensue, with fibrovascular membrane formation and traction retinal detachment.

HIV retinopathy
Untreated advanced HIV disease is associated with a retinopathy characterized by multiple cotton-wool spots, usually without accompanying haemorrhages (Fig. O.19). It is asymptomatic and usually clears with
Effective antiretroviral therapy. It may occasionally be difficult to distinguish this from early cytomegalovirus (CMV) retinitis, which typically occurs in patients with CD4+ lymphocyte counts below 50 per mm³. CMV retinitis causes irreversible retinal damage and is characterized by large perivascular exudates and haemorrhages (Fig. O.20). Retinitis due to toxoplasmosis has also been described (Fig. O.21).

**ORTHOPNOEA**

Alex West

(See also DYSPNOEA, p. 144.)

A patient is said to have orthopnoea if he experiences breathlessness when lying down. Most patients suffering breathlessness at rest are more comfortable in the sitting position, but a clear history of intense breathlessness in the recumbent position is particularly characteristic of left ventricular failure, mitral stenosis and other conditions that cause pulmonary venous hypertension. The paroxysmal nocturnal dyspnoea of patients with left ventricular failure is at least partly related to posture in bed. Episodic breathlessness at night is also a common feature of asthma and, as asthma symptoms can be problematic at night, this needs to be carefully distinguished from the true orthopnoea of pulmonary oedema.

**OTORRHOEA**

Michael Gleeson

Otorrhoea (aural discharge) may arise from the external auditory meatus, middle ear or, occasionally, intracranial or adjacent structures. It is an unpleasant and worrying symptom for the patient. At the outset, it is important to note the character of the discharge, whether it is watery or mucoid, continuous or intermittent, its colour, its quantity and whether it is inoffensive or offensive. Some discharges are very subtle, and all require a very thorough inspection of the ear with a microscope and microsuction. The causes of discharge are listed in Box O.9.

**EXTERNAL AUDITORY MEATUS**

Wax (cerumen) is so variable in colour and consistency that it can be mistaken for something else. Most often just present in small flakes or boluses, it can change as a result of infection or water contamination into a light yellow, semi-liquid discharge. Wax is nature's disinfectant. It provides the first barrier against infection, and dissolves squamous epithelium shed from the tympanic membrane. Wax is transported laterally by an inherent pattern of epithelial movement and, in normal circumstances, should not need to be removed. In some situations, it may become impacted in the deep meatus or contaminated with water; in these circumstances,
microsuction or syringing is appropriate. *Keratosis obturans* is a condition where wax and desquamated squamous epithelium form an adherent mass in the bony meatus, sometimes even causing bony erosion. This condition may be associated with bronchiectasis and sinusitis as part of the *immotile cilia syndrome*.

**Otitis externa**

This is a very common disease and is sometimes part of a generalized dermatitis, for example eczema or psoriasis. Some patients may precipitate the condition by the use of hairsprays, perfumes and even eardrops to which they are allergic. Indeed, even some wax solvents can cause enough irritation to create the condition.

The most common cause of otitis externa, however, is a breakdown in the migratory mechanism of the skin of the deep meatus referred to above. Elsewhere in the body where skin divides, movement, clothes, friction and washing all serve to dispose of the dead skin. The external meatus is not exposed to any of these factors, so a different mechanism exists to dispose of the dead epithelium. The viable epithelium in the area has the property of migrating dead epithelium from the deep meatus to the wax glands in the outer part of the ear canal. If, for any reason, this migratory process breaks down, dead epithelium collects in the deep meatus. The entrapped skin acts as an excellent culture medium for bacteria, particularly if it is wet. Secondary bacterial infection takes place rapidly, giving rise to symptoms of pain and discharge. Treatment consists of thorough aural toilet and antibiotic drops. It may take some time for the mechanisms within the external ear to return to normal. Throughout this time, the patient should be careful not to scratch the ear and to avoid further water contamination.

*Fungal otitis externa* will not respond to the usual antibiotic eardrops, and it may even be caused by their overuse. It has a distinct appearance so can be recognized immediately without recourse to fungal cultures. In these patients, the ear canal is filled with moist, whitish debris that looks and suction like damp blotting paper. In *Aspergillus niger* infection, black spots are seen amid the fungal debris. The other common causative fungus is *Candida albicans*.

Viral otitis externa exists in two forms, each of which can cause a type of otitis externa. Reactivation of *herpes zoster* causes excruciatingly severe pain and may sometimes be accompanied by facial paralysis (*Ramsay Hunt syndrome*) (Fig. O.22). The other type is *myringitis bullosa haemorrhagica*, which is often seen in conjunction with the flu virus. Blood blisters form on the drum and lead to a serosanguineous discharge and bouts of very severe pain. Both of these conditions cause pain that is significantly more severe than the clinician would expect from the physical signs alone.

In many cases, it is not possible, because of oedema, to see the tympanic membrane. It is most important to differentiate between simple otitis externa and otitis externa secondary to otitis media as soon as possible. One useful differentiating feature is that the pain of external otitis is precipitated or exacerbated by movement of the pinna, while in otitis media it is pressure above and behind the ear that is painful. If, in addition to the characteristic pain, the patient has good hearing and a positive Rinne test, even though the meatus may be almost completely occluded, the condition is most unlikely to be otitis media.

*Malignant otitis externa* is a poor term as it wrongly implies a neoplastic process. Unfortunately, this term has become enshrined in otological practice and is unlikely to be changed. It refers to a *Pseudomonas* or fungal infection of the tissues of the skull base. This infection gains access by a breach of the skin in the external auditory canal at the junction of the bony and cartilaginous parts. Malignant otitis externa
is a very severe and sometimes fatal infection that tends to develop in those who are either elderly or immunocompromised for any reason – for example, with diabetes, chemotherapy or AIDS. Scant discharge precedes the spreading infection that presents with consummate, unremitting pain together with progressive cranial nerve palsies. In late cases, gross destruction of the skull base takes place.

More correctly, malignant otitis externa should refer to patients with carcinoma of the external auditory canal. This too is extremely painful, is almost always misdiagnosed, and is underestimated for months until the discharge becomes blood-stained and facial palsy supervenes. Fortunately, it is a very uncommon form of squamous cell carcinoma as it has a poor prognosis (Fig. O.23).

Rarely, a salivary fistula involves the cartilaginous meatus and may follow injury that involves the ear, the temporomandibular joint and the parotid gland. Discharge can also be caused by a first branchial arch sinus that classically communicates with the bony cartilaginous junction, and may pass through the parotid and intermingle with branches of the facial nerve (Fig. O.24).

OTORRHOEA

Acute otitis media gives rise to discharge once the tympanic membrane has ruptured. Until then, the patient suffers from pain, increasing deafness and malaise. The pain subsides as soon as the pus is released. Initially, the discharge is blood-stained, but it soon becomes frankly mucopurulent and often profuse, pouring down the cheek and soiling the patient’s pillow. This continues for some days until the infection comes under control, as a result of either antibiotic therapy or natural immunity. If the eardrum is inspected carefully, the perforation can be seen, through which a pulsating discharge escapes. The pus is usually inoffensive. Acute otitis media is caused by either viral or bacterial infection. The organisms most commonly implicated are Streptococcus, pneumococcus and Haemophilus.

Chronic suppurative otitis media is classified as either tubo-tympanic or attico-antral. The hallmark of tubo-tympanic otitis media is a central perforation, a defect in the pars tensa of the drum. Few ears with persistent perforations are entirely dry. Most are moist, with episodes of increased and sometimes profuse discharge during acute exacerbations. In some cases, it may be possible to close the perforation by surgical means, thereby eradicating the discharge. In others, this is not possible, and the discharge must be kept to a minimum by strict water precautions and intermittent courses of antibiotics. Continuous infection in the middle ear cleft stimulates hyperplasia of the mucosal lining, which ultimately prolapses through the perforation as a polyp.
The fundamental lesion of attico-antral chronic otitis media disease is an attic retraction pocket, with or without cholesteatoma, or ingrowth of skin from a marginal perforation. In some cases – particularly children – polyps develop adjacent to the attic defect. The precise cause of these pathological processes is not known. Cholesteatoma, a nidus or collection of moist and often infected epithelium, releases enzymes that help to destroy bone and erode the temporal bone. As a result, the cholesteatoma can extend into the mastoid, disrupt the ossicular chain, and destroy the bony labyrinth and facial nerve. Given time, the cholesteatoma eventually abuts the dura of the middle and posterior cranial fossas, from where infection can spread intracranially and either meningitis or brain abscess will develop. This disease can be extremely serious, and the cholesteatoma should be eradicated by mastoid surgery.

Primary malignant neoplasia of the middle ear is extremely rare, but it usually presents with a blood-stained, mucopurulent discharge arising from granular tissue that has destroyed the tympanic membrane. The diagnosis can only be made by biopsy. Malignant tumours arising in the middle ear cleft include squamous cell carcinoma, metastatic tumours and eosinophilic granuloma (histiocytosis X).

Cerebrospinal fluid (CSF) can escape from the ear and appear as a watery discharge. Although spontaneous CSF leaks are possible, they would have to develop in a patient with a perforated eardrum in order to drain from the ear rather than the Eustachian tube. Most patients with CSF otorrhoea have a clear history of severe trauma, sufficient to have fractured the temporal bone.

OVERACTIVITY

Andrew Hodgkiss

The causes of overactivity may be classified as shown in Box O.10.

Disorders of movement may frequently provide valuable clues about a diagnosis, especially when a patient is confused, mute or otherwise unable to give a clear account of symptoms. Disorders of movement include overactivity, underactivity (see p. 697) and abnormal involuntary movements.

Overactivity describes behaviour where there is an increase in physical activity, over-talkativeness and sometimes aggressiveness. The subject may make exaggerated gestures and facial expressions, and at interview have difficulty in sitting still, feeling impelled to move about the room, their attention easily distracted by external stimuli. This type of overactivity typically occurs in mania, when it is associated with elevated or irritable mood, pressure of speech and flight of ideas, grandiose plans and, in severe forms, grandiose or paranoid delusions. Sleep is disturbed, and patients may remain awake and active all night, with apparently undiminished energy levels next morning. Libido and appetite may both be increased. Such a typical presentation of a manic illness presents few diagnostic problems, especially if there is a history of previous mood swings or a family history of affective disorder.

Restless overactivity may also be drug-induced or a feature of organic brain disease. Both acute and chronic organic confusional states (see CONFUSION, p. 102) may present with irritability, restlessness and excitement, and this can be particularly marked in states of drug intoxication and withdrawal from alcohol and sedatives. In the early stages of Alzheimer’s-type dementia, a state may develop resembling mania, and the diagnosis should be made from the history of increasing memory impairment. Neurosyphilis could also present in this way, and associated signs such as Argyll Robertson pupils, peripheral neuropathy and evidence of dementia combined with serological tests will confirm the diagnosis. Patients with temporal lobe epilepsy may rarely develop post-ictal confusional states with overactivity, irritability or senseless aggression. These episodes are usually brief and in the context of a

<table>
<thead>
<tr>
<th>Box O.10 Causes of overactivity</th>
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<tbody>
<tr>
<td><strong>Common causes</strong></td>
</tr>
<tr>
<td>• Mania</td>
</tr>
<tr>
<td>• Organic brain disease (see CONFUSION, p. 102)</td>
</tr>
<tr>
<td>• Delirium</td>
</tr>
<tr>
<td>• Agitated depression</td>
</tr>
<tr>
<td>• Anxiety</td>
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<tr>
<td>• Schizophrenia</td>
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<tr>
<td>• Hyperthyroidism</td>
</tr>
<tr>
<td>• Hyperparathyroidism</td>
</tr>
<tr>
<td>• Drugs</td>
</tr>
<tr>
<td>• Caffeine intoxication</td>
</tr>
<tr>
<td>• Withdrawal from: sedatives, hypnotics, anxiolytics or alcohol</td>
</tr>
<tr>
<td>• Abuse of hallucinogens (LSD), amphetamine, cocaine or phencyclidine</td>
</tr>
<tr>
<td>• Akathisia and restless leg syndrome</td>
</tr>
<tr>
<td>• Attention-deficit hyperactivity disorder in children</td>
</tr>
<tr>
<td><strong>Less common causes</strong></td>
</tr>
<tr>
<td>• Post head injury (in children)</td>
</tr>
<tr>
<td>• Anorexia nervosa</td>
</tr>
<tr>
<td>• Temporal lobe epilepsy (post-ictal confusional state)</td>
</tr>
<tr>
<td>• Neurosyphilis</td>
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</table>

...
known history of epilepsy. Akathisia is a particular form of restlessness found frequently in Parkinson’s disease and as a side effect of antipsychotic medication. Subjects are unable to remain still through a subjective sense of unease and restlessness. They have an urge to get up to move their feet and legs or rock the body. Restless legs may keep them awake at night and be troublesome to a partner.

Overactivity may be a main presenting feature of agitated depression, when the subject’s complaint of worry is reflected in their appearance, thought content and behaviour. The person will appear restless, tense and fidgety, constantly seeking reassurance, because of feelings of guilt about the past and uncertainty about the future. Agitation may also be a principal feature of an anxiety disorder, and in patients with schizophrenia. In all these psychiatric conditions and in physical illnesses including hyperthyroidism and hyperparathyroidism, agitation may be associated with signs of increased autonomic activity including sweating, tachycardia, palpitations, shallow breathing and gastrointestinal disturbance.

Overactivity without significant mood disturbance can be a feature of anorexia nervosa. These patients will exercise relentlessly and methodically in order to keep their weight low. The exercise may take the form of housework or regular training in a gym, or various sports.

Hyperkinesis is the most common and most disruptive sequela of head injury in children. Features include restlessness and impulsive disobedience at home and at school, sometimes with explosive outbursts of anger. A similar pattern of hyperactivity and resistance to discipline has been observed in children with epilepsy and after encephalitis lethargica, although epidemic encephalitis has fortunately now largely disappeared. Both organic and psychogenic factors are involved, and the pre-traumatic personality of the child and the family setting may be as important as the severity or nature of the head injury.

The attention-deficit hyperactivity disorder (ADHD) in children is viewed as a developmental abnormality presenting around the time the child enters school or earlier, with inappropriate degrees of inattention, impulsiveness and hyperactivity. The disorder is manifest in all situations, including at home, at school and in play activity. There may be inappropriate running about in the classroom, fidgetiness and over-talkativeness. These children tend to be academic underachievers with low self-esteem, low mood and temper outbursts. There is an increased likelihood of ‘soft’ neurological signs, including poor coordination. Enuresis and encopresis are more common. Children may show features of conduct disorder, the impairment of social and school functioning persisting throughout childhood.
PAIN, PSYCHOGENIC

Andrew Hodgkiss

Pain is the most common complaint brought to a doctor, and it is probably the most complex of subjective experiences. All pain has sensory, emotional and cognitive dimensions that require management. For example, there is a large literature to show that patients who are well informed about surgical procedures preoperatively suffer less pain and anxiety postoperatively and require less analgesia. Thus, the ‘psychological aspects’ of all acute and chronic pains need managing. However, it is the large number of patients with chronic, medically unexplained pain that will be focused upon here.

A large proportion of chronic pain remains medically unexplained. For example, the findings of half of all diagnostic laparoscopies for chronic pelvic pain look normal. Many patients disabled by headache, facial pain and back pain have repeatedly normal investigation results. Are these chronic pains of psychological origin?

Two routes of psychogenesis of pain have been described throughout the twentieth century: a hysterical mechanism (the conversion of a psychological conflict into a meaning-laden somatic symptom) and pain as a ‘depressive equivalent’. These two rather different psychogeneses have tended to be lumped together under the term ‘psychogenic pain’ or pain disorder. A good example of a hysterical mechanism is the chronic pelvic pain of the female patient sexually abused in childhood. The pain can be interpreted as a result of the repressed memories of this trauma. Pain as a depressive equivalent rests on the idea that some patients suffer in a non-localized way and report low mood to doctors, while others, lacking words for emotional states (alexithymic patients), localize their suffering within the space of their bodies, and report regional bodily pain instead. These patients often have a positive family history of depressive illness.

The treatment of patients with pain disorder is now well established. They are offered pain management programmes with multidisciplinary components including psychologists and occupational therapists. A cognitive–behavioural approach is usually adopted, with pain behaviour being left unrewarded, and habits of thought when in pain being examined and corrected. Opiate dependence, which often comes to complicate the chronic pain, is tackled. In addition, antidepressant medication, particularly with tricyclics, is valuable. Despite the hysterical mechanism underlying a proportion of these pains, it was Freud’s experience, and that of others since, that psychoanalytic treatment is less successful with pain than with other hysterical somatic symptoms.

It is best to reserve the use of the term ‘psychogenic pain’ for those patients in whom positive evidence of psychogenesis is evident. The rest are best left in the category of ‘medically unexplained symptoms’. There is growing interest in and understanding of how this large number of patients should be managed.

A number of psychiatric disorders can present with pains, for example muscular tension pains in anxiety disorders (especially non-cardiac chest pain), a gripping abdominal pain in depression or hallucinatory pain in schizophrenia. Pain is a popular symptom choice for those with factitious disorders and malingering.

Finally, it is important to distinguish psychogenic pain from neuropathic pain. Confusion sometimes arises clinically because both respond to tricyclic antidepressants. However, neuropathic pain is medically explicable and has a characteristic distribution and nature, for example the burning sensation in the soles of the feet in the sensory peripheral neuropathy associated with alcohol misuse.

PALLOR

Mark Kinirons

Pallor is a very subjective sign indicating a reduction in skin or mucous membrane colouring. It usually refers to a generalized reduction rather than to localized depigmentation, as in vitiligo.

Pallor may be a congenital characteristic resulting in an individual always looking pale even when healthy. Anaemia, for any reason, is the most common cause of pallor. Pallor of the conjunctivae or mucous membranes is a very unreliable indication of a reduced haemoglobin, except when there is severe anaemia. Pallor may be accentuated in hypopituitarism when, in addition to the anaemia, there is a reduction in skin pigmentation.

A reduction in blood flow to the skin will result in acute pallor. This can arise either due to hypotension, as in shock, or due to hypothermia or low cardiac output states. Similarly, profound vasoconstriction (as seen in phaeochromocytoma) will also lead to pallor and is the mechanism by which emotional shock or stress can be responsible for pallor.
Any increase in the thickness of the skin, particularly the epidermis, will tend to obscure the haemoglobin in the capillaries, resulting in pallor. Skin thickness is increased in acromegaly and myxoedema.

**PANIC**

**Andrew Hodgkiss**

Panic is an extreme form of anxiety characterized by (i) severe, acute, brief attacks, and (ii) the feeling of loss of personal control. The symptoms are predominantly the somatic features of anxiety (especially cardiovascular) and hyperventilation, so that a primarily physical presentation is common. The emotions experienced are stark and distressing: terror, fear of dying, collapsing, going mad or losing control. Attacks may be triggered by specific stimuli or be unpredictable, and life frequently becomes dominated by the apprehension of the acute distress and helplessness engendered. Panic attacks are common, occurring at some time in 10 per cent of the adult population and to a persistently disabling degree in one in five of those affected. The causes of panic are listed in Box P.1.

There are a number of physical conditions that enter the differential diagnosis. Hyperthyroidism and phaeochromocytoma can mimic closely while, particularly in a diabetic or partial gastrectomy patient, the possibility of hypoglycaemia should be excluded. Other episodic medical conditions that can be confused with somatic presentations of panic attacks are migraine, Ménière’s disease, temporal lobe epilepsy and carcinoid syndrome. Not uncommonly, patients with these conditions develop genuine panic attacks as an emotional reaction to the uncertainty they experience: they may begin to report symptom differences between attacks, which can confound the physician unless he or she is aware of the possibility of a superimposed panic disorder.

In addition to the symptom content, the brief duration of panic attacks (usually lasting a few minutes) is a pointer in distinguishing panic from most physical disorders.

Panic attacks are frequently predominant features in the setting of other psychological conditions. In phobias, panic is often experienced when exposed to the phobic stimulus, but anticipating or even thinking about exposure may trigger panic. Panic attacks occur, and may occasionally be the presenting complaint in, depressive illness, or they may develop subsequently during recovery; episodes occurring regularly early in the day are a clue, but there are usually readily identifiable features of depressive change evident in the history and mental state. Hyperventilation may be involved in the development of panic attacks and may also be exacerbated by them.

Panic attacks are induced by stimulants, both illicit drugs like amphetamine and cocaine and the socially acceptable caffeine – a note of coffee consumption is worth taking. Drug withdrawal also precipitates episodes; perhaps most commonly implicated are the benzodiazepines and alcohol.

Finally, the relationship between anxiety and panic is disputed. It is evident that patients who suffer from generalized ‘free-floating’ anxiety are likely to experience panic attacks of varying degree and frequency, while patients having panic attacks get appreciably anxious about these alarming experiences and commonly develop a secondary chronic anxiety state or agoraphobia. Panic disorder is now considered a separate diagnostic entity with putative differences in inheritance, pathophysiology, natural history and treatment responses compared with generalized anxiety disorder.

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**Box P.1  Causes of panic**

### Most common
- Normal
- Stress-related
- Anxiety disorder
- Panic disorder
- Agoraphobia with panic attacks

### Less common
- Drugs
  - Stimulants
  - Hallucinogens
- Drug withdrawal
- Physical disorders
  - Hyperthyroidism
  - Hypoglycaemia
- Phaeochromocytoma
- Carcinoid syndrome
- Asthma
- Ménière’s disease
- Temporal lobe epilepsy
- Migraine
- Psychiatric disorders
  - Depression
  - Chronic hyperventilation syndrome
  - Post-traumatic stress disorder

### Rare
- As for anxiety (see p. 30)
WHITE OR PEARL Y PAPULES

Milia, small firm white lesions, are commonly seen on the face, and are commonly multiple. Strictly they are not papules but minute keratin containing cysts. They may follow blister formation (e.g. porphyria cutanea tarda and epidermolysis bullosa) or appear after an episode of sunburn.

Keratosis pilaris is a common condition of children and young adults, and consists of rough, firm, white papules, approximately 1 mm in diameter. It is found over the lateral upper arm and the anterior thighs above the knees; the buttocks are occasionally involved. The papules give the skin a nutmeg grater-like feeling, and they are occasionally surrounded by an inflammatory halo. They are caused by keratin plugs in the ostia of hair follicles. The development of keratosis pilaris is determined by a dominant gene, and the condition is closely related to ichthyosis and atopic dermatitis. Rarely, it is a manifestation of vitamin A deficiency (phrynoderma).

Molluscum contagiosum is a common viral infection (Fig. P1), occurring especially in children and young adults. It is readily transmitted from person to person, and in addition individual lesions may be autoinoculated. Typical lesions present as pearly white or skin-coloured papules. They have a smooth shiny surface with central umbilication, from which a cheesy material may be expressed, and the typical inclusion bodies may be demonstrated under the microscope.

In children, especially atopic subjects, lesions may be widespread on the face and limbs; in adults, they are more common around the genitals and may be acquired through sexual contact. After a period of some months, the body mounts an immune reaction, and the lesions become acutely inflamed and then resolve spontaneously, sometimes leaving a minute scar.

Syringomas are common benign lesions, frequently multiple, arising from the sweat duct apparatus, and they usually occur on the upper cheeks just below the eyes as smooth flesh coloured oval papules 2–4 mm in length. Lesions accumulate over the years and must be differentiated from xanthelasmas, which are yellower and larger and fewer in number and may be associated with hyperlipidaemia. In the rare disorder lipoid proteinosis, pearly translucent white papules occur widely on mucous membranes and skin, particularly the eyelid margins.

SKIN-COLOURED PAPULES

Viral warts are a common cause of skin-coloured papules and occur particularly over the dorsa of fingers and on the soles. Flat plane warts can also occur on the backs of hands and may form subtle tan papules over the chin and cheeks, especially in young women. In older subjects, viral warts are uncommon, and small multiple keratotic papules on the backs of the hands or scalp are more likely to be solar (actinic) keratoses.
The diagnosis of a solitary viral wart in an older subject must always be treated with suspicion, as an early squamous cell carcinoma may present in this way. Clustered subcutaneous papules formed into an annular configuration are suggestive of granuloma annulare. They are seen most commonly on the fingers, backs of hands, elbows or ankles, and resolve spontaneously over a number of years (Fig. P.2).

**BROWN PAPULES**

*Basal cell papillomas* (seborrhoeic warts) – the most common benign tumour of the elderly – may grow to form large brown plaques, but they begin as light brown tiny verrucous papules scattered widely on the chest and back.

*Genital warts* are usually small, brown, flat-topped and smooth with an angular outline. This is in contrast to the *condylomata lata* seen in secondary syphilis, where extensive crops of soft, skin-coloured or pink pedunculated papules or nodules, covered with tiny moist papillary projections, occur on the genitalia, scrotum and perianal region (and also between the toes and at the angles of the mouth). The serology will be positive.

An *acrochordon* (skin tag) is a tiny, brown, pedunculated soft papule with a narrow neck found in the flexures of patients from middle-age onwards. Lesions may be numerous around the neck, axillae and groins. They are more profuse in the obese, and are particularly marked in *acanthosis nigricans*. In *Darier’s disease*, the papules are brownish, perifollicular and covered by a brown–grey crust. The papules may be grouped or accumulate into sheets, and the sites of predilection are the seborrhoeic areas of the scalp, behind the ears, and in the nasolabial folds, mid-back and interscapular regions. They may be crusty on the scalp, and vegetating in the intertrigenous areas. Palmar pits and notching of the nails are associated features that may assist in the clinical diagnosis; if in doubt, the histology is characteristic. The disease often flares during the summer months, and it is dominantly inherited.

Dermatosis papulosa nigra is the rather grand and cumbersome title of tiny common jet black papules over the cheeks of black subjects. In *lichen amyloidosi*s multiple, closely spaced, uniform, rounded, hard papules that may be red, brown or skin-coloured occur over the anterior shins, or rarely more widely. The lesions are intensely pruritic, resistant to treatment and not associated with systemic amyloidosis. Even rarer are the brown/skin-coloured papules crowded around the nose seen in *tuberose sclerosis*, the so-called *adenoma sebaceum of Pringle* (Fig. P.3). Other cutaneous manifestations are shagreen patches, ashy leaf hypopigmented macules and periungual fibromas. There may be associated neurological, cardiac and renal problems.

**VIOLOCEOUS PAPULES**

Flat-topped shiny polygonal papules are the hallmark of *lichen planus*. The colour is characteristically lilac–pink. The papules are smooth and not crusted, and they may be crossed by tiny white lines known as Wickham’s striae. They are intensively irritating and may appear in areas of skin damage, such as a scratch, this being termed the Koebner or isomorphic phenomenon. The eruption is usually symmetrical, particularly on the flexural aspects of wrists and forearms, anterior and inner aspects of calves and ankles, and over the upper abdomen and lumbar region. In Caucasians,
a delicate, white, lace-like pattern is common on the buccal mucosa, and this is an important diagnostic sign. Similar lesions may be present on other mucous membranes (medial aspect of labia majora or the glans penis). Lesions may be simulated by a lichenoid drug eruption; mepacrine, oral hypoglycaemics, beta-blockers and gold have all been implicated. Lichen planus resolves spontaneously after a year or two, leaving macular post-inflammatory pigmentation at the previously affected sites.

**BLACK PAPULES**
A solitary uniform sharply marginated blue–black papule is highly suggestive of a blue naevus. The lesions remain unchanged over many years, and malignant transformation is vanishingly rare. The sudden appearance of multiple, uniform black papules may indicate satellite spread of a malignant melanoma. In multiple idiopathic haemorrhagic sarcoma of Kaposi, dark purplish papules appear usually over the feet and ankles of elderly patients of Jewish or Italian lineage. Gradually, the lesions become elevated to form multiple papules and later soft nodules, plaques and angiomatous tumours. They may ulcerate, bleed and crust. A more generalized, aggressive and metastasizing form of the disease is endemic in East Africa and is seen in association with HIV infection.

**YELLOW PAPULES**
Xanthelasmas are found on the upper cheeks and eyelids in middle-aged men and women. Investigation should exclude hyperlipidaemia, coronary artery disease, diabetes, myxoedema or primary biliary cirrhosis. The sudden appearance of numerous yellow papules over the buttocks and elsewhere indicates eruptive xanthoma, a condition in which abnormalities of lipid metabolism are invariably. Naevoxanthoendothelioma (juvenile xanthogranuloma) occurs as rounded, yellow, firm papules or nodules over the extensor surfaces within a few weeks of birth. They involute spontaneously in 1–3 years and are not associated with underlying lipid abnormalities. The skin of the neck and axillae may be covered in sheets of yellowish papules in pseudoxanthoma elasticum. This is inherited as an autosomal recessive condition and is a generalized defect of support tissue. There may be associated tears in Bruch’s membrane in the eye, and aneurysms of small and large arteries.

Tiny yellow papules on the face and forehead of older people can be caused by sebaceous hyperplasia; the lesions are usually flat-topped, with an erythematous halo and a tiny central depression. They must be differentiated from molluscum contagiosum, which is more rapidly growing and numerous, and early basal cell carcinoma, which is more translucent in colour and is associated with telangiectasia and slow growth with ulceration. Discrete tiny yellow papules 1–2 mm in diameter can be due to ectopic sebaceous glands visible on the lips or mucous membranes, penis and vulva (Fordyce’s disease).

**RED PAPULES**
Several of the more common skin diseases have papular elements or a papular stage in their evolution (e.g. psoriasis, acne, dermatitis and chickenpox). Psoriasis (see SCALY ERUPTIONS, p. 599) may present as myriads of tiny, flat, pink papules scattered widely over the body and limbs; this cutaneous reaction pattern is known as guttate psoriasis, and may follow a streptococcal infection. This eruption occurs in adolescence and may clear over a few weeks, but papules can amalgamate into the more characteristic plaques. Pityriasis lichenoides chronica produces lesions somewhat like guttate psoriasis but with more confluent papules. Each is surmounted by a mica-like scale that can be detached en masse. This occurs in young individuals, and it may persist for months or even years; an acute form (pityriasis lichenoides acuta) exists in which chickenpox-like inflammatory lesions predominate, although mixed forms also exist. The eruption of pityriasis rosea occurs over the T-shirt area (trunk, upper arms and tops of thighs) in the typical ‘Christmas tree’ arrangement of dozens of oval, salmon-coloured macules. These scale from within outwards, leaving a characteristic collarette of scale near their margins. The eruption is often preceded by a ‘herald patch’, which is a larger (but otherwise typical) lesion, often situated on the trunk. Sometimes, especially in children with dark skins, the lesions may be more papular, not unlike lichen planus or papular syphilide. Purpuric papules are the hallmark of a cutaneous vasculitis, and all cases must be investigated to exclude an associated systemic component (Fig. P.4). If sudden in onset, meningococcal septicaemia must be excluded; if there is any doubt as to the possibility of this diagnosis, antibiotic therapy must be initiated at once.

Comedones are the cardinal signs of acne vulgaris, and these may co-exist with papules, pustules, cysts and scars. Consequently, the lesions of acne vulgaris are follicular, and this differentiates them from the papules of rosacea, which are not. In rosacea, comedones are absent, and scarring does not occur. Dermatitis and eczema often have a papular phase before the lesions progress to vesicles and weeping.
When eczema becomes chronic, the skin becomes thickened and leathery, with exaggerated skin markings (lichenification). In *lichen simplex chronicus*, which has some eczematous features, large oval patches of skin-coloured lichenified papules are seen on the typical areas (nape of neck, ankles and perineum). The condition is most common in women between the ages of 30 and 50, and is severely pruritic. A variant of this is *nodular prurigo*.

In a *papular syphilide*, the papules are shiny or scaly, but the colour is of a copper-red hue. The palms, soles and forehead are particularly affected, and there is no pruritus. Other features of secondary syphilis may be present, including fever, lymphadenopathy, patchy alopecia and ‘snail-track’ erosions in the mouth and genital areas. *Campbell de Morgan spots* are cherry-red and smooth-surfaced. They can be seen in most elderly people on the trunk and proximal parts of the limbs. Similar (but slightly smaller and scaly) lesions are *angiokeratoma*; these are common on the scrotum of old patients (angiokeratoma of Fordyce) and rare in the bathing trunk distribution in Anderson-Fabry disease (alpha galactosidase deficiency), a rare inborn error of metabolism with cutaneous, renal, vascular and neurological manifestations.

**PARANOID DISORDERS**

Andrew Hodgkiss

In everyday language, paranoid ideas are considered synonymous with unpleasant persecutory ideas. However, in a stricter sense, the word ‘paranoid’ means a morbid disturbance of the relationship between self and others. This can encompass themes of persecution, jealousy, love, guilt, grandiosity, self-reference and so on.

The majority of paranoid beliefs do involve ideas of persecution, taking the form of delusions or overvalued ideas, and these have limited diagnostic specificity. Ideas of persecution develop in normal individuals deprived of sleep or at times of great stress. Quite commonly, they may be transient phenomena in immigrants living in a new and strange environment without home supports. Other predisposing factors include deafness, especially when accompanied by failing physical powers and loneliness in the elderly, although, curiously, blindness does not appear to be such a strong predisposing factor as deafness.

Paranoid ideas may become more evident as age advances in individuals who have long-standing personality traits of undue sensitivity and wariness in relationships. Such people may have few friends and social supports. There is a tendency for those with paranoid illnesses to be living alone. Simple persecutory delusions may occur in the early stages of a dementia or be due to a brain tumour or drug abuse, particularly with stimulants such as amphetamines.

Paranoid delusions developing in middle life in the absence of other psychotic phenomena used to be dubbed ‘paranoia’ but are now known as the delusional disorders. These conditions were very well delineated in twentieth-century French psychiatric classification, and many still bear the name of the psychiatrist who first described them. The French were always certain that these syndromes arose from the personality and autobiography of the individual, and that they were very different from schizophrenia.

*Erotomania* (de Clérambault’s syndrome) describes a delusional disorder, usually found in women. These women develop an unshakeable conviction that a man, usually an authority figure, is in love with them, and they may develop elaborate erotic delusions and cause considerable social disruption in attempting to come to the attention of the object of their feelings. Erotomaniac delusions may occur as part of a schizophrenic or affective psychosis too.

In *morbid jealousy*, a spouse, usually the male, will become unreasonably suspicious concerning their
partner’s fidelity. It takes the form of an overvalued idea when the subject can be temporarily reasoned out of their conviction, or it may be held with delusional intensity when, despite evidence to the contrary, the spouse remains intensely jealous and preoccupied. Morbid jealousy is sometimes associated with alcoholism, schizophrenia and affective illness; it may occur in psychopathic persons, and sometimes in the absence of any other psychiatric pathology. The symptom is a cause of domestic violence and not infrequently homicide.

Hypochondriacal delusions similarly are a common presenting symptom in schizophrenia and in psychotic depressive illness, but they may be the only feature of a delusional disorder, in which the subject will be convinced of infestation of the skin by insects, or their bowel by worms, that their body gives off a bad smell, or that part of their body is not working (intestines, stomach or brain) or is not there at all (nihilistic delusions). The patient’s preoccupation may centre around the teeth, gums, mouth and face and, until the delusional quality of the belief is recognized, it may lead to excessive medical and surgical referrals and requests for investigations. These delusions may be accompanied by social phobias if the abnormal belief includes a conviction that part of the body is deformed, or to sexual difficulties if the delusions relate to the sexual organs.

In Capgras syndrome, the patient becomes convinced that someone close to them, often a spouse, has been replaced by a double who resembles the spouse in appearance but is really an imposter. This is most commonly a symptom in a schizophrenic illness, but, again, it may stand alone as a delusional disorder. An unusual feature of paranoid psychosis is that, in rare instances, paranoid beliefs are contagious and are taken up by the healthy partner of the psychotic patient. This particularly occurs if the partner is relatively isolated and dependent on the sick person, who often has paranoid schizophrenia. A folie à deux situation may persist for years, maintaining stability in the partnership, each reinforcing the other’s beliefs. Separation leads to complete loss by the normal partner of the paranoid ideas.

PARAPLEGIA

David Werring & Mark Kinirons

Paralysis of both legs may occur with damage to any part of the neural pathways between the motor cortex supplying the legs and the muscles of the legs themselves. In practice, paraplegia most commonly results from disorders of the spinal cord, spinal roots or peripheral nerves. If the onset is acute, it can be difficult to distinguish spinal from neuropathic paralysis, because the initial spinal shock results in the abolition of reflexes and is associated with flaccidity. In acute spinal cord disorders with damage to the corticospinal tracts, the paralysis affects all muscles below a given level, and it is often associated with sensory loss below the same level. Frequently, there is accompanying bladder and bowel paresis.

In peripheral nerve disorders, motor weakness tends to involve the distal muscles of the legs rather than the proximal ones, and sphincter function is usually spared. Sensory loss, if present, is also more prominent in the distal parts of the limbs.

For clinical purposes, it is helpful to separate the acute paraplegias from those of gradual onset. The former should always receive urgent attention, otherwise neurological damage, in particular loss of sphincter control, may become permanent, even in surgically remediable conditions. It should be noted that, in several disorders, for example multiple sclerosis and spinal cord compression, the onset of paraplegia may be either sudden or gradual.

SUDDEN ONSET

The most common cause of acute paraplegia is spinal cord trauma, usually combined with fracture dislocation of the spine; spontaneous haematomyelia and spinal cord infarction resulting from thrombosis of one of the arteries supplying the spinal cord (usually the anterior spinal artery) are less common causes.

Post-infectious myelitis, acute demyelination, epidural abscess or tumour with spinal cord compression, and epidural or subdural haemorrhage from bleeding disorders or anticoagulant therapy may cause acute paraplegia, often associated with severe pain. Other causes include loss of power in the legs due to acute poliomyelitis or acute idiopathic post-infectious polyneuritis.

GRADUAL ONSET

In adults, (i) multiple sclerosis, (ii) subacute combined degeneration, (iii) spinal cord compression from tumours, (iv) cervical discs, (v) cervical spondylosis, (vi) epidural abscess, (vii) motor neurone disease, (viii) syringomyelia and (ix) degenerative disease of the lateral columns represent the most frequently encountered forms of spinal paraplegia of gradual onset. Parasagittal tumours bilaterally compressing the motor cortex representing both legs are another rare and readily overlooked cause of paraplegia. Finally, the possibility that paraplegia is a manifestation of hysterical illness should be borne in mind.
Pelvic Pain

Tony Hollingworth

All women will at some time experience pelvic pain associated with events such as menstruation, ovulation or sexual intercourse. Although only a few women seek medical advice for such pain, it remains the most common reason for laparoscopic examination in the UK. Visceral pain in the abdominal and pelvic organs is transmitted by the autonomic nervous system in the T10–L1 distribution to the lower abdomen and is usually a dull sensation with vague localisation. The viscera are not very sensitive to thermal or tactile sensation, and they are poorly localized in the cerebral cortex. Stimuli that produce pain are as follows:

• Distension and/or contraction of a hollow organ.
• Rapid stretching of the capsule of a solid organ
• Any irritation of the parietal peritoneum, for example by blood or pus
• Tissue ischaemia or necrosis, as may happen in torsion of an ovarian cyst
• Neuritis secondary to any inflammatory, neoplastic or fibrotic processes in adjacent organs.

The differential diagnosis of pain in the pelvis can be divided into:

• Acute, when the patient is ill and requires resuscitation with intravenous fluids for hypovolaemia, sepsis or dehydration. There may be a need for urgent surgery.
• Subacute, in which the onset of the pain is sudden but does not cause the patient to be severely ill. The diagnosis may need to be established, although it may just require observation over 24–48 hours. If the pain does not improve or the diagnosis is in doubt, laparoscopy may be warranted.
• Chronic in nature and possibly the result of longstanding pathology. Symptoms, clinical findings and findings at laparoscopy show poor correlation, with up to 50 per cent of patients who undergo laparoscopy showing no abnormality.

Acute and Subacute Pain

The differential diagnosis for this type of pain can be classified in the following way. Investigations and treatment may be similar, but severity can vary, and the need for any surgical intervention will be a clinical decision. In all these cases, it is important to consider a possible physiological cause or a pregnancy-related condition.

• Physiological: Menstruation or ovulation. Some women habitually experience some dull pain in the midline or in one or other iliac fossa at the time of ovulation, usually 14 days before the next period (Mittelschmerz). Occasionally, slight vaginal bleeding accompanies the pain. The timing of the pain and the absence of any abnormal pelvic findings usually make the diagnosis clear. Pain is not an unusual feature with menstruation, and it usually implies that the woman is ovulating and that there is an increase in local prostaglandins within the uterus at that time.
• Pregnancy-related: It is always important to remember to consider pregnancy in any woman during her reproductive years when she presents with acute pelvic pain. The severity of pain will dictate the management. Miscarriage may be related to bleeding, and pain usually reflects the inevitable nature of the miscarriage; on examination the cervical os will be open. Ectopic pregnancy, which is still a major cause of maternal mortality, may present with varying symptoms depending on the gestation. Although this problem can be treated conservatively with methotrexate, the vast majority of cases will need some form of operative intervention. Other problems in pregnancy include fibroid degeneration, which may be very acute and need hospital admission. Apart from pain, it can also cause the pregnant uterus to become very irritable.
• Ovarian: Ovarian causes of acute pain usually reflect cyst accidents in the form of torsion, rupture or haemorrhage (Fig. P5). The degree of peritonism will dictate the management.
• Infection: Pelvic inflammatory disease (PID) occurs in the sexually active. It is usually bilateral in origin, and it is associated with a low-grade pyrexia, tachycardia and discharge. If not adequately treated
initially, tubo-ovarian abscesses (Fig. P.6) can occur, which may need surgical drainage.

- **Endometriosis:** This condition may cause both acute and chronic pelvic pain, and diagnosis is usually confirmed by laparoscopy.
- **Neoplasia:** This includes fibroids, ovarian cysts and malignant disease within the genital tract. Apart from the conditions mentioned above, it is unusual for malignancies to give acute pain.
- **Urinary tract:** Other non-gynaecological causes include urinary tract infection, retention and renal stones.
- **Gastrointestinal:** The degree of acuteness may vary, and the pathology would vary depending on the age of the patient. Diagnoses include appendicitis, gastroenteritis, constipation, diverticular disease, inflammatory bowel disease, acute hernial accidents, volvulus, mesenteric infarction and malignancy.

### Chronic Pelvic Pain
Any acute cause of pain can lead on to a chronic picture. A history of pelvic pain is associated with an increased number of sexual partners, and an increased incidence of psychosexual trauma as a child. The investigations are similar, may lead to laparoscopy and indeed, in some cases, may result in hysterectomy and pelvic clearance. Although there is a place for this with endometriosis and chronic PID, it will not result in cure for all women if there is no obvious pathology.

- **Adhesions:** these may be found in up to 20 per cent of patients with chronic pain (Fig. P.7), although they may not be the cause of the pain.
- **Residual ovary syndrome:** this occurs in women who have had a hysterectomy and is associated with pain, dyspareunia and a fixed tender ovary at the vaginal vault. Treatment may be by hormonal suppression to establish the diagnosis or surgical removal or release.
- **Endometriosis:** in this condition, there is abnormal implantation of endometrium outside the uterine cavity. It may cause dysmenorrhoea, dyspareunia, menstrual upset, pelvic pain and infertility. Diagnosis is usually by laparoscopy. Adenomyosis is a variant of this condition when endometrium invades the myometrium, and this causes pain with the period and a very tender uterus. The surgical option is dependent on symptoms.
- **Chronic PID:** this is a consequence of acute infection and leads to damage and consequent pain and menstrual upset.
- **Irritable bowel syndrome:** this condition is often confused with a gynaecological cause of lower abdominal pain.
- **Pelvic congestion:** the pain is dull and aching with occasional sharp exacerbations and is associated

### Routine Investigations
These may vary from patient to patient, but a full blood count and pregnancy test may be sufficient depending on the history. If infection is considered, swabs should be taken from the vaginal vault and endocervix to exclude *Chlamydia*, and from the urethra and rectum if gonorrhoea is a possibility. A mid-stream specimen of urine may exclude a urinary infection. Plain abdominal X-rays may be useful if bowel problems are an option. An intravenous pyelogram or KUB (kidney, ureter and bladder) ultrasound scan may exclude a calculus. The mainstay of investigation is the ultrasound scan of the pelvis and, when in doubt, laparoscopy may be undertaken. However, many that are undertaken do not reveal any obvious pathology.
with dilated veins in the broad ligament and uterus. There may be local tenderness. These women are usually in their reproductive years, and may be nulliparous. Diagnosis can be made by venography, laparoscopy or ultrasound (Fig. P.8), and treatment is medically with progestogens.

- **Psychological:** depression, anxiety or somatization may predispose or contribute to pain in the adult. There may be a past history of abuse which may contribute to the problem. This may be an avenue to explore if no other diagnosis can be identified.


**PELVIS, SWELLING IN**

Tony Hollingworth

Swellings that arise from the pelvis can be considered under their anatomical origins. A number of structures may appear to be pelvic when their original site of origin is really abdominal. Ultrasound scanning has improved the detection of lesions that are not necessarily palpable without a vaginal or rectal examination. The background to the swellings can be simply described by the five Fs, namely fat, fluid, faeces, flatus and fetus. Careful history-taking, clinical examination and appropriate imaging (ultrasound scan and magnetic resonance imaging) should be able to establish the diagnosis. Anatomically, swellings will arise from the following.

**BLADDER**
- Simple distension or retention
- Transitional cell carcinoma (see HAEMATURIA, p. 241)

The most common difficulty that arises in the diagnosis of pelvic swellings is to differentiate between a distended bladder, pregnant uterus, ovarian cyst and uterine fibro-myoma. The distended bladder is the easiest to exclude by the passage of a catheter to establish the diagnosis. Neglect of this simple procedure has led to the abdomen being opened.

**VAGINA**
- Haematocolpos
- Hydrocolpos

Distension of the vagina by menstrual fluid is not likely to be mistaken for anything else, if only on account of the absolute closure of the atretic membrane that causes it. This condition is often referred to as an ‘imperforate hymen’. This is not correct because the atresia is at a higher level in the vagina than the hymen, which is always perforate. Haematocolpos is practically the only central tumour occurring between the rectum and the bladder reaching from the hymen to the pelvic brim. It usually presents in girls between the ages of 16 and 17 years, with acute retention of urine due to the fact that the swelling fills the pelvis, and the distended bladder in front is forced upward into the abdomen. Primary amenorrhoea is present, although monthly menstrual cycle symptoms without loss of blood may have taken place. Two swellings are present: the tender distended bladder in the lower abdomen which can reach as high as the umbilicus, and the distended vagina filled with menstrual fluid in the pelvis. The uterus can usually be felt like a cork movable upon its upper extremity. The lower pole of the haematocolpos presents a blue-coloured swelling at the vulva (see Fig. M.13). A similar swelling may be found on rare occasions in newborn girl babies: the vagina is filled with a milky fluid – hydrocolpos.

**UTERUS**
- Pregnancy-related, either normal or abnormal, with or without associated tumours of the uterus or ovary
- Non-pregnancy-related, the most common of which include fibroids (leiomyomas), haematometra and pyometra (blood or pus in the uterine cavity), endometrial carcinoma and, more rarely, uterine sarcoma or chorionic carcinoma

History is of great value in differentiating these cases, with amenorrhoea being usual with pregnancy problems, menorrhagia and pressure effects common with fibroids. Bleeding can occur in early pregnancy at the time when the periods would be due, or as a result of threatened miscarriage. Fibroids may cause menorrhagia depending on their location within the uterus (Fig. P9). Other pathologies tend to be linked with age. Haematometra may be related to cervical stenosis following treatment to the cervix for precancer; pyometra may occur in postmenopausal women and suggest malignancy. Endometrial carcinomas may
present with menstrual upset, but most present in the postmenopausal age group with bleeding or discharge. Palpation can be difficult to differentiate a gravid uterus in the early stages of pregnancy as the uterus may fluctuate like a cyst, a softened fibroid may do the same, or a tense ovarian cyst may feel so hard as to be mistaken for a fibroid. It can be difficult to detect the fetal heart by sonicaid in the first trimester. If the pedicle of the mass can be felt definitely attached to one uterine cornuum, it is strong presumptive evidence of an ovarian tumour. When small tumours are in question, the first point to establish is whether the tumour can be separated from the uterus bimanually. If it can, it is unlikely to be either a fibromyoma of the uterus or a normal uterine pregnancy. This distinction can only be made by careful bimanual examination, and in some cases this will require considerable skill. A pedunculated fibroid is, of course, extraterine and may have the same anatomical arrangement to the uterus as an ovarian tumour. If it is a fibroid that has undergone cystic change, the physical signs are identical to those of an ovarian cyst.

Early pregnancy in a retroverted uterus may give rise to diagnostic difficulties but it should be remembered that the soft, cystic fundus is felt through the posterior fonsix, that the cervix looks down the vagina or forwards to the symphysis, and that the posterior mass is continuous with the cervix. If the retroverted uterus is associated with bladder distension, the picture is usually clear enough. The history of urinary retention followed by constant dribbling of urine (distension with overflow), amenorrhoea, other signs of pregnancy, the presence of two tumours – one in front, tense, tender and elastic; the other behind, soft and cystic – and finally the passage of a catheter will settle the question. The diagnosis of solid ovarian tumours is not always possible as the pedicle is often short, and the tumour is then so close to the uterus that the two cannot be separated and can be mistaken for fibroids.

In the case of definite uterine tumours, age is important. The diagnosis of cancer of the endometrium is usually in postmenopausal women and the size of the uterus may be small. The diagnosis should always be confirmed by microscopic examination of curetted fragments. Fibroids are only likely to be mistaken for malignant growths when they produce constant bleeding as a result of extrusion, degeneration and sloughing. The rapid growth of a fibroid is more likely to be the result of degenerative changes than to the developing of a sarcoma. Growth of a fibroid after the menopause, however, should make one consider the need to exclude sarcomatous change within it.

**OVARY**

- Benign – includes cysts and fibromas
- Malignant – primary origin in the form of epithelial tumours (85 per cent), sex cord tumours (6 per cent), germ cell tumours (2 per cent) and uncommonly sarcomas or lymphomas. Secondaries (6 per cent) originate from the gut, breast, lung and thyroid

Large tumours arising from the pelvis are often not difficult to differentiate from one another, bearing in mind the differential diagnosis includes ovarian tumours; uterine fibroids and pregnancy are the most commonly occurring conditions. It cannot be repeated too often that amenorrhoea plus a pelvic mass needs pregnancy to be excluded. Menorrhagia usually suggests uterine fibroids. Exceptions to these general statements are uncommon, and mistakes in diagnosis will occur but seldom if these are borne in mind. Ascites has to be differentiated from ovarian cysts and, in general, gives dullness in the flanks on percussion, with resonance over an area somewhere around the umbilicus. Ovarian cysts, which can be bilateral, give dullness over the front of the abdomen, with resonance in the flanks. When ascites exists along with ovarian tumours, the free fluid may be so large in volume that the tumour cannot be felt. As a rule, however, it can be felt on pressing through the fluid, and the omentum may be ballotable as an omental cake due to secondary spread. Ascites with
an ovarian tumour does not always mean malignancy. Some fluid may also accompany fibroma of the ovary, or a simple ovarian cyst with a twisted pedicle. Ovarian fibromas may be accompanied by a large amount of ascites and bilateral pleural effusions (Meigs’ syndrome). Some of the ascitic fluid should routinely be sent for cytology.

When pregnancy is associated with a tumour, the diagnosis may be difficult. This does not lie in the recognition of the pregnancy; amenorrhoea, breast changes, fetal movements and the fetal heart will usually make that clear enough. It lies in deciding the nature, or even the presence, of a tumour along with the pregnant uterus. In the early months, when the presence of two tumours can be demonstrated, the diagnosis is easier, but in the later months with the increasing size of the abdomen, the swellings may appear to merge into one another and obscure the picture. Fibroids are likely to soften and degenerate during pregnancy, so that they are liable to be mistaken for ovarian cysts.

In the case of ovarian tumours, it is often impossible to be sure of the exact nature of the growth until it has been removed and examined histologically. Because of this doubt, there should be no undue delay in the removal of an ovarian tumour larger than the size of an orange or one that is growing. Small follicular cysts may be left as they are harmless and eventually disappear. Fixation of the growth in the pelvis, obvious ascites, unilateral oedema of the leg, emaciation of the patient, abdominal pain and a rapid growth in size of the abdomen point to malignancy.

As a rule of thumb, a cyst less than 5 cm in diameter may resolve without any action except repeat scanning after two or three normal periods. If the cyst is larger than 5 cm, it will probably need to be formally removed. Very large cysts tend to be either benign or borderline malignant on histology (Fig. P10). The largest cyst removed in the UK weighed 63 kg; the world’s largest was removed in 1905 in the US and weighed approximately 145 kg. The ovary should not be palpable in a postmenopausal woman, and any ovarian cyst in these women should be considered malignant until proved otherwise.

**FALLOPIAN TUBES**

- Pregnancy-related, tubal gestation or progressive extraterine pregnancy
- Inflammatory – salpingitis, which may lead to a hydrosalpinx or pyosalpinx
- Malignant, carcinoma of the Fallopian tube being very uncommon

With small tumours confined to the pelvis or rising only a little above the brim, diagnosis is often difficult. In practice, however, extraterine gestation and its resulting blood tumour stand out pre-eminently as a swelling. Before rupture or abortion has occurred, a tubal gestation is essentially a small tumour in one posterolateral corner of the pelvis, attached to the uterus, indefinite in consistency, remarkably tender and perhaps – although not always – associated with amenorrhoea of short duration and acute attacks of pain in the pelvis. Definite signs of pregnancy may not be present, but a pregnancy test will be positive.
It may be mistaken for a chronic salpingo-oophoritis, a small cystic ovary, a small pedunculated fibroid or a small ovarian dermoid. The differential diagnosis may be difficult, but attacks of pain not associated with menstruation are not likely to occur in any of the above conditions; the pains are usually the result of overdistension and stretching of the tube from haemorrhage into its wall or lumen around the fertilized ovum.

When tubal abortion, or tubal rupture, has occurred, the signs of internal bleeding, accompanied by sudden pain and collapse, with haemorrhage from the uterus or the passage of a decidual cast, usually make an unmistakable picture. Intraperitoneal haemorrhage is more commonly severe and copious with tubal rupture than with tubal abortion. If the patient recovers from the initial bleeding, the clinical picture may be that of a retro-uterine or peritubal haematocoele. The uterus is pushed forwards and upwards against the posterior fornix pubis, and the mass of blood clot can be felt posteriorly bulging the posterior fornix and also the anterior wall of the rectum. Vaginal examination can be very tender. Tubal miscarriage is most likely to be mistaken for an ordinary intrauterine miscarriage, but the presence of a tender mass on one side of the uterus, with a closed cervix and a negative ultrasound scan, and the absence of uterine contractions or the extrusion of any products of conception, should make the case clear. Pain is much more severe, but external bleeding is much less in an extrauterine pregnancy compared to a miscarriage.

The essential point in diagnosing an ectopic pregnancy is to approach every woman of child-bearing age who complains of irregular bleeding and abdominal pain with that possibility in mind. With the improved resolution of vaginal scanning and the development of early pregnancy assessment units, many ectopic pregnancies are picked up early and can be treated medically with methotrexate. The rest are usually dealt with laparoscopically and it is seldom that a laparotomy is required these days. No two cases are alike, and there are more exceptions to the rule in the symptomatology of this condition than in any other. It must be emphasized that, although maternal death is not common in the UK, ectopic pregnancy remains a major cause of it.

Progressive extrauterine gestation is a rare occurrence and is the consequence of continued growth of an embryo after a partial separation from the tube as a result of rupture or extrusion from the fimbriated end (abortion). The continued enlargement of a mass beside the uterus with amenorrhoea and progressive signs of pregnancy are the most characteristic points. Abdominal pain in late pregnancy is a characteristic feature.

The uterus may be felt in the pelvis separate from the fetal sac. The diagnosis, however, is difficult, because there is always some effused blood, which obscures the outlines of the uterus, and makes it appear to be a part of the pelvic mass. The fetus is often situated high above the pelvis, and it tends to lie transversely facing downwards. A radiograph reveals the fetus adopting a position that is characteristically odd, the spine hyperextended or acutely flexed, and the head and limbs at unusual angles to the trunk. If, on a lateral view, radiography shows fetal parts overlapping the maternal spine, the pregnancy must be extrauterine. Ultrasonography will establish the absence of an intrauterine gestation and also the size of the uterus, which never exceeds that of a 5 months’ gestation even in the presence of a full-term extrauterine pregnancy, and the cervix does not soften to the same degree. In those cases where the fetus lies in the front of the false sac, it will feel very superficial owing to the absence of uterine wall in front of it, and between it and the examining hand.

The swellings due to salpingo-oophoritis are usually easy to distinguish. They form fixed tender masses in the pelvis, seldom of any definite shape, but occasionally presenting the characteristic retort shape, with its narrow end near the uterus, which the tube assumes when distended with fluid. The history is usually that of an acute illness at some period, with pain in the pelvis, a rise of temperature and peritoneal irritation. It is preceded, as a rule, by uterine discharge and menorrhagia. This inflammatory disturbance can be associated with long periods of infertility, owing to the sealing up of the tubes. In the chronic state, pelvic pain, congestive dysmenorrhoea, dyspareunia, vaginal discharge, menorrhagia and infertility are all possible symptoms. The signs of suppuration, pyrexia and leucocytosis, wasting and daily sweating are usually absent, and the pus in the tubes is sterile.

A large pelvic abscess may accompany salpingo-oophoritis, or may occur alone without infection of the tubes, as we occasionally see in puerperal septic infections. When it does occur, it is of course peritoneal; it fixes the uterus in a central position, bulges into the posterior fornix and rectum, tends to rupture into the rectum, before which there is a copious discharge of mucus per anum, is acute in onset, and is accompanied by signs of local peritonitis. A swinging temperature, leucocytosis, sweats and the symptoms of fever are present, all suddenly improving when the abscess discharges itself. It is likely to be confounded with pelvic cellulitis, in which the uterus is fixed in a laterally displaced position. This swelling bulges one
patients tend to display altered bowel problems. Gynaecological tumours, as is diverticulitis. These felt. Bowel cancer is more common than the common interval between it and the iliac crest can usually be tumour when an ovarian cyst is present, and some ovarian pedicle. There is usually a definite fluctuating acute onset may be similar to that of torsion of an rarely fluctuates unless there is a large abscess. The adherent to the iliac fossa. The lump is ill defined and anterior superior spine of the ilium, and apparently inflammation is, however, in close relation to the an ovarian pedicle. The swelling due to appendix may be mistaken for such a condition as torsion of an pelvically.

BONE
Growth of the pelvic bones is very rare and usually cartilaginous or sarcomatous. This is only likely to be mistaken for adherent inflammatory masses due to salpingo-oophoritis. They will be found to be continuous with the bones forming the pelvis and, when growing from the sacrum, may have the rectum in front of them; all other pelvic tumours have the rectum behind them. In most cases of this nature, the uterus and adnexae can be palpated bimanually and shown to be free from disease and unconnected with the mass. When complicated by the presence of a pregnant uterus, their true nature may be difficult to determine unless examination reveals that they are absolutely fixed and continuous with the bones of the pelvis.

OTHER STRUCTURES
Many of these lesions are not primarily pelvic, but they are included in the list because they are liable to be mistaken for pelvic tumours. Thus, renal, splenic or pancreatic tumours may reach the pelvic brim, but the history ought to show that they have grown down from above, not up from below. Renal swellings may be associated with urinary changes, or absence of urinary secretion on the affected side as detected by the cystoscope or an intravenous pyelogram. Malformations of the genital tract are associated with developmental abnormalities of the renal tract. It is not uncommon to find a solitary pelvic kidney in patients with congenital absence of the vagina and uterus. Splenic enlargements may be associated with blood changes. Pancreatic cysts are the least likely to be mistaken for pelvic swellings, but they have been difficult to distinguish from ovarian tumours with long pedicles.

PENILE LESIONS
Ben Challacombe
Sores on the penis may be present on the thin mucous covering of the glans or prepuce, or on the cutaneous surface of the body of the penis; they are more common in the former situation. Penile examination should always involve retraction of the prepuce/foreskin, and lesions in this region drain to the inguinal nodes, which should always be assessed clinically. Ulceration in the neighbourhood of the glans penis may be due to:

- Infective lesions
  - Balanitis
  - Genital herpes
  - Chancroid
  - Granuloma venereum (inguinale)
– Lymphogranuloma inguinale (venereum)
– Chancre
• Papilloma
• Lichen sclerosis et atrophicus (balanitis xerotica obliterans)
• Psoriasis
• Premalignant lesions
• Squamous cell carcinoma (SCC)
• Kaposi’s sarcoma

**Balanitis**
This usually occurs in uncircumcised men when inflammatory processes have been allowed to continue beneath the prepuce. This results in ulceration and excoriation of the mucous membrane covering the glans penis or lining the prepuce, accompanied by a purulent discharge. The prepuce often becomes swollen and oedematous, preventing retraction, so that a condition of phimosis occurs or, if retraction has taken place, the analogous state of paraphimosis, due to a tight band almost strangulating the end of the penis. Simple balanitis may be due to *Candida albicans* infections (particularly in those with diabetes), but it may also result from acute gonorrhoeal urethritis, or an underlying syphilitic or soft chancre. Balanitis cincnata is part of Reiter’s syndrome, occurring in association with urethritis, arthritis, conjunctivitis (often slight and transient) and buccal ulceration. In Behçet’s syndrome, too, ulcerative penile and scrotal lesions may occur in association with buccal ulcers. With an acute urethritis, there will be a history of infection and pain along the course of the urethra during micturition (see DYSURIA, p. 154); the intracellular gonococcus may be identified in a Gram-stained smear of the discharge. Zoon’s balanitis is an inflammatory condition producing a shiny erythematous plaque on the glans or prepuce, and often both these sites where they appose each other with the foreskin forward causing ‘kissing lesions’. Treatment is usually by circumcision.

With a chancroid (soft sore) due to bacterial infection, consecutive sores may appear about the orifice of the prepuce, while the inguinal nodes are much more likely to be inflamed or to suppurate than with simple balanitis. A syphilitic chancre, occurring 4 weeks after infection, can be obscured by a phimosis, but it can usually be felt distinctly under the skin and causes a comparatively small amount of discharge, while the inguinal nodes become enlarged but do not suppurate.

**Herpes Genitalis**
Herpes may attack the genital organs as part of a herpes zoster or more commonly herpes simplex type II infection, an important sexually transmitted disease. The disease begins as a patch of erythema on the inner surface of the prepuce or on the glans penis, followed by vesicles and pustules. Swabs must be sent at presentation, and herpes simplex virus often recurs despite antiviral therapy. Soft chancres are usually deeper, with marked edges; their bases are sloughing, and they are usually accompanied by a bubo (enlarged inguinal lymph node, which may suppurate), which is exceptional with herpes. A syphilitic chancre is usually single, indurated and raised, and is accompanied by the typical multiple, discrete nodes in the inguinal region. It should be remembered that syphilis may become inoculated upon a herpetic patch, or that herpetic may appear in an area already infected with syphilis.

**Chancroids or Soft Sores**
Chancroid is a bacterial sexually transmitted infection caused by the fastidious Gram-negative streptobacillus *Haemophilus ducreyi*. It is a disease found primarily in developing countries, associated with commercial sex workers and their clientele. Soft sores or chancroids of the penis have a short incubation period, with vesicles occurring in 2 days. These break down rapidly to form a rounded or oval ulcer with undermined edges and a yellowish sloughing base. The ulcers appear usually on the mucous surface of the glans, frenulum or corona, and are multiple. They may cause rapid destruction of tissue, perforating the frenulum or spreading over the surface of the glans. Chancroid must be differentiated from a syphilitic chancre, and serum from the edge of the lesion should be examined for *Haemophilus ducreyi* (Ducrey’s bacillus), as well as by dark-ground illumination for *Treponema pallidum*. Co-infection with chancroids and syphilis is not uncommon. The chancroids are multiple and are accompanied by a thin, purulent discharge and by a painful swelling of the inguinal nodes, which have a tendency to suppurate. A syphilitic chancre is single, raised and indurated, has little discharge and is accompanied by enlarged but firm and indolent nodes in both inguinal regions; the incubation period of a syphilitic chancre is from 21 to 28 days. The multiple ulcerations caused by herpes are more superficial. Uncircumcised men are at three times greater risk than circumcised men for contracting chancroid from an infected partner. Chancroid is a risk factor for contracting HIV, due to the ecological association or shared risk of exposure, and biologically facilitated transmission of one infection by the other due to ulceration. Treatment is with a single oral dose (2 tablets) of azithromycin or a single intramuscular dose of ceftriaxone or oral erythromycin for 7 days.
GRANULOMA INGUINALE
Granuloma inguinale (granuloma venereum) is caused by the bacterium *Calymmatobacterium granulomatis*. The disease is commonly found in tropical and subtropical areas such as south-east India, Guyana and New Guinea, and is spread by vaginal or anal intercourse. It is a chronic granulomatous ulceration that may affect the perineum and the inguinal regions, as well as the penis. The painless lesions on the penis start as a papule, which appears after a few days’ or weeks’ incubation period, and breaks down to form a superficial ulcer. Syphilis is the main differential diagnosis. Examination of the discharge shows intracellular capsulated Gram-negative rods known as Donovan bodies (*Donovania granulomatis*). Lymph nodes are not involved, and treatment is with erythromycin, streptomycin or tetracycline antibiotics.

LYMPHOGRANULOMA VENEREUM
Lymphogranuloma venereum (lymphogranuloma inguinale) or tropical bubo is another sexually transmitted disease that is common in the tropics but also seen in patients with HIV. It is a chronic condition characterized by initial small, painless ulcers on the penis, with marked glandular enlargement in the groins and severe constitutional disturbances. The organism travels from the site of inoculation down lymphatic channels to multiply within mononuclear phagocytes of the lymph nodes it passes, which then tend to break down and form sinuses. The lesion on the penis appears, after an incubation period of about a week, as a vesicle, papule or ulcer, and it tends to disappear by the time the lymphatic nodes are enlarged. It is due to a filter-passing organism, *Chlamydia trachomatis*, which can be diagnosed by serology and polymerase chain reaction, or a biopsy of the primary lesion or lymph node. Rectal stricture and effusions into joints are other lesions caused by this disease. Treatment is with antibiotics such as tetracycline, doxycycline and erythromycin.

SYPHILITIC CHANCRE
Chancre, the initial lesion of syphilis, generally appears on the penis about 25 days after infection, and it is most common in the region of the frenulum or coronary sulcus. A chancre appears as a reddened patch, which becomes raised above the surface of the mucous membrane, with distinctly indurated margins. The central part breaks down into an ulcer (Fig. P.11), discharging a thin, purulent fluid, and at the same time the inguinal nodes of both sides become palpable, slightly enlarged, but discrete, and with no tendency to suppurate. The chancre increases but slowly in size, or may occasionally become smaller without any treatment, and after a further lapse of 4–6 weeks, if the condition remains untreated, the typical secondary symptoms make their appearance – namely, a roseolar rash on the chest, abdomen, face and thighs, general adenitis, and mucous patches about the faucial pillars and tonsils, accompanied by low pyrexia. The diagnosis of the primary lesion of syphilis frequently presents no difficulties, the indurated character of the sore, the date of its appearance after infection, and the presence of firm, indurated nodes in the inguinal region being distinctive. If the character of the sore is not distinctive, it is necessary to differentiate it from other lesions of the penis. A careful search must be made by dark-ground illumination for the *Treponema pallidum* in the serum expressed from the sore; negative serological reactions in the early stage of the disease are not reliable. If the sore is syphilitic, the secondary manifestations of the disease will follow. Diagnosis of a chancre is difficult when it is hidden beneath a phimosed prepuce. There is a purulent and foul discharge from beneath the oedematous and swollen prepuce; the inguinal nodes are enlarged from the associated sepsis. If any doubt exists as to whether an indurated subpreputial area is an SCC or a syphilitic sore, circumcision should be performed under anaesthesia, the ulceration inspected, biopsied and sent for histological analysis, and some serum expressed from the ulcer examined on a dark stage for *Treponema pallidum*.

LICHEN SCLEROSIS ET ATROPHICUS
This is a chronic dermatitis that is known as balanitis xerotica obliterans when present on the penis. It causes itchy, flat-topped, white papules that coalesce to form white patches. It is a progressive condition that results in phimosis and urethral meatal stenosis. It is also a
premalignant condition, but progression to SCC is rare. Patients should be followed up and initially treated with short courses of topical steroids. If the condition is severe or progressing, circumcision or urethral meatal dilatation may be required.

**PSORIASIS**
Psoriatic lesions may appear guttate (raindrop-shaped), circinate (rings) or geographical. Genital psoriasis may involve the glans, and this can be treated with topical emollients or short courses of topical low-dose steroid creams.

**PREMALIGNANT PENILE CONDITIONS**
There are a number of other premalignant cutaneous lesions of the penis, and any chronic red or pale lesion on the glans or prepuce should be reviewed after topical treatments, and a biopsy performed if persistent. Such conditions include:
- Lichen sclerosis/balanitis xerotica obliterans: steroids if mild, or circumcision to diagnose
- Cutaneous horn: a solid skin overgrowth, treated by wide local excision
- Pseudoepitheliomatous micaceous and keratotic balanitis: hyperkeratotic growths requiring excision
- Leucoplakia: solitary or whitish plaques involving the meatus. Excise and follow up
- Erythroplasia of Queyrat: carcinoma-in-situ of the glans, prepuce or penile shaft. It is a red, velvety, circumscribed and painless lesion. Excise and treat with laser ablation, topical 5-fluorouracil or radiotherapy
- Bowen's disease: carcinoma-in-situ of the perineal skin. Widely excise, and treat with laser or cryoablation
- Buschke–Lowenstein tumour: verrucous carcinoma or giant condyloma. An aggressive locally invasive tumour of the glans. Excise widely

**SQUAMOUS CELL CARCINOMA (SCC)**
Squamous cell carcinoma is the most common malignant growth of the penis, accounting for 95 per cent of penile malignancies (Fig. P.12). Risk factors include age (sixth to eighth decades), premalignant lesions (see above), a foreskin, smoking and human papillomavirus types 16 and 18. A solid, non-tender mass or ulcer arises from the inner aspect of the prepuce (21 per cent), or from the mucous membrane of the glans (48 per cent), coronal sulcus or shaft. The lesion is often a small, raised ulcer with friable, irregular edges, and it frequently occurs on the site of previous ulceration or long-standing irritation; it is unknown where circumcision has been performed in infancy. It grows to involve the corpora cavernosa, urethra and eventually the perineum. It is often associated with enlargement of the inguinal lymph nodes, which may be due to septic infection or malignant infiltration. Metastases go to the superficial and deep inguinal nodes, and then to the iliac and obturator nodes before disseminating. It may be confused with a chancre, but the friable, irregular edges, the liability to bleed and the gradual progressive increase in size, in an elderly patient, should give rise to suspicion of malignant disease.

Management involves a high index of suspicion in non-healing ulcers despite treatment. Biopsy is confirmatory and graded as low, intermediate or high. Patients often present late due to embarrassment, personal neglect, fear or ignorance. Treatment should be carried out in large specialist centres and involves total or partial penectomy, preferably with penis-sparing techniques. High-grade and high-stage disease may require inguinal lymphadenectomy and inguinal sentinel node biopsy, and advanced disease is treated with chemo-radiotherapy.

**PAPILLOMAS**
Genital warts (condylomata acuminata) occur on the glans and contiguous surface of the prepuce and are most frequently found on the corona. They are related to human papillomavirus type 16 and 18 infection.
They are usually multiple, and are distinguished from SCC by the absence of induration in the base. Treatment is with podophyllin 5 per cent, and a biopsy often rules out SCC. Bowenoid papules are thought to be due to human papillomavirus infection and resemble carcinoma-in-situ, but they have a benign course and should be biopsied.

**KAPOSI’S SARCOMA**
This is a reticuloendothelial tumour that is now increasingly common due to its occurrence in immunocompromised men with HIV/AIDS. It begins as a raised, painful, bleeding, violaceous papule, but it can be a blue ulcer. It is slow-growing and often solitary but diffuse on occasion. Lesions may cause urethral obstruction, and treatment is with laser therapy, cryoablation, radiotherapy or intralesional chemotherapy.

**TUBERCULOUS ULCERATION**
Tuberculous ulceration of the penis is rare and generally associated with advanced tuberculous infiltration elsewhere. Tuberculous ulcers are usually shallow, with thin overhanging edges, and they are painful and multiple. The diagnosis is clinched by discovering tubercle bacilli in films made from the discharge.

**INJURY**
Injury is an uncommon cause of a penile sore but may follow a zipper injury, a penile bite, which needs to be treated with broad-spectrum antibiotics, or a sporting injury.

**PENIS, PAIN IN**

Ben Challacombe

Pain in the penis is a symptom that occurs not only in association with lesions of the penis or urethra, but also as a referred pain from disease of the prostate, bladder or kidney. Penile pain may be present either during or immediately after micturition, or it may be entirely independent of the act. If pain is felt only during micturition, there is probably some inflammatory lesion of the urethra or prostate; if it occurs immediately after the flow of urine, it suggests some lesion in the urinary bladder; and pain present quite apart from micturition may be due to various diseases of the penis, bladder, ureter or kidney.

The term ‘pain’, too, is a relative quantity, varying with the nervous susceptibility of the patient, as what is pain in one may be merely discomfort in another, so that the patient’s description may have to be discounted to a certain extent by the clinician.

**CAUSES OF PAIN IN THE PENIS EXPERIENCED DURING MICTURITION**
- Diseases of the urethra
  - Acute inflammation, gonorrhoeal/non-specific urethritis (NSU), chemical or other
  - The passage or impaction of a calculus
  - Stricture of the urethra
  - Injury to the urethra
  - Foreign body in the urethra
- Diseases of the prostate
  - Acute prostatitis
  - Prostatic abscess
  - Prostatic carcinoma
- Diseases of the bladder
  - Acute cystitis
  - Bladder calculus
  - Pedunculated bladder carcinoma

**Diseases of the urethra**
The most common cause of pain in the penis during micturition is acute inflammation of the urethra, often gonorrhoeal, but it may result from other organisms, and this is particularly common following catheterization. Non-specific urethritis, a common sexually transmitted infection, is diagnosed when gonorrhoea and other bacterial infections have been excluded. Frequently, this is due to *Chlamydia trachomatis* or *Trichomonas vaginalis* (see URETHRAL DISCHARGE, p. 722). In the earliest stages of an acute urethritis, before any marked urethral discharge is apparent, there is usually a sense of smarting or tingling in the terminal urethra, more marked as the discharge increases, when it is of a burning or scalding character. The pain during micturition occurs within a few days of sexual intercourse and is frequently the earliest symptom of urethral infection; a purulent discharge from the urethra is usually present later.

The passage of a calculus through the urethra causes a sharp, cutting pain along the urethra, the cause of which is apparent when the calculus is voided. A stone may pass into the urethra during micturition and become stuck at a narrowing, usually at the membranous portion or at the distal end, when a sudden sharp pain is felt, and the flow of urine is partially or completely stopped. Occasionally, a calculus may remain in the urethra behind a stricture in the bulb. The whole length of the urethra should be examined by passing the finger along its course, when a stone may be actually felt; or the calculus may be seen through a cystoscope or identified on a plain radiograph.

*Urethral stricture* occasionally causes pain during micturition, especially if the calibre is small, and if there is infection or ulceration of the urethral mucosa.
behind the stricture, but as a rule stricture causes little pain. The common symptoms are a gradual increasing difficulty in micturition, a poor stream and dribbling of urine from the meatus after the stream has terminated; the diagnosis will be confirmed by a flat and prolonged flow rate, leading to direct observation of the urethra through a flexible or rigid cystoscope.

*Injury to the urethra* may cause pain during micturition. The urethra may be injured by a fall on the perineum, by a kick or blow, or by the faulty or careless passage of instruments; it may also be injured or lacerated in association with a fracture of the pelvis. The urethra may be merely bruised, lacerated on one aspect or completely ruptured. If it is lacerated by direct injury, blood usually appears at the external urinary meatus, together with a contusion in the perineum or along the course of the urethra; any attempt at micturition causes pain in the penis, while urine may or may not be expelled from the meatus, depending upon the extent of the injury, or may be extravasated into the perineal or scrotal tissues (Fig. P.13). A suprapubic catheter will usually be needed to relieve obstruction. An ascending urethrogram will delineate the site and severity of the urethral injury.

A *foreign body* in the urethra may cause considerable pain. In some cases, the history will be clear – for instance, the end of a catheter may have broken off within the urethra – but in others, no history of the insertion of a foreign body into the urethra will be forthcoming. Urethroscopy will show the foreign body; various articles have been found in the urethra, such as a wire, string, wood and hairpins.

**Diseases of the prostate**

*Acute prostatitis* and *prostatic abscess* both give rise to pain during micturition, in addition to increased frequency and difficulty during the act. Both are usually sequelae of an acute urethritis but they may follow prostatic biopsy and an abscess may result in undulating fluctuations in temperature. The diagnosis of the two conditions is made by rectal examination and, if an abscess is present, a softer, fluctuant and acutely tender area in the inflamed gland can usually be detected.

Benign prostatic hyperplasia (BPH) generally does not cause penile pain during micturition, but pain in the penis is occasionally present during micturition in cases of prostatic carcinoma, owing to the direct infiltration of the urethral mucous membrane. On rectal examination, prostate cancer feels hard and irregular and occasionally fixed (T3/4), in contrast to the elastic, smooth feel of BPH. A serum prostate-specific antigen test with or without transrectal ultrasound-guided biopsy may be indicated.

**Diseases of the bladder**

Diseases of the bladder may cause penile pain during micturition in certain circumstances, although it is more common to find that pain in vesical disease follows the completion of micturition. In acute cystitis, penile pain is present throughout micturition, due to the intense congestion of the trigone. The other symptoms of acute cystitis, namely suprapubic pain, pyrexia, increased frequency of micturition and the presence of pus and blood in the urine, suggest the diagnosis.

Pain during micturition in other vesical lesions is caused whenever there is sudden obstruction to the normal flow of urine by the impaction of something against the internal urethral orifice. This may occur with a small calculus, a foreign body or a pedunculated tumour, accompanied by a shooting pain in the urethra, while after an interval of a few seconds the stream may be re-established. With vesical calculus, the urine may be normal or may contain pus and blood if the bladder has become infected; there is penile pain after micturition, and the stone will be seen both on imaging (plain X-ray of the pelvis, ultrasound or MRI) and with a cystoscope.

**Penile pain following micturition**

This symptom is common to many lesions of the urinary bladder, more especially those in which there is ulceration or infiltration of the basal areas. The particular pain felt by the patient is described as a sharp prickling or tingling at the terminal part of the penis on cessation of micturition, lasting some minutes and causing a desire to squeeze the glans.
The possible causes of pain in the penis following micturition are:

- **Vesical**
  - Calculus
  - Tuberculosis
  - Tumour
  - Acute cystitis
  - Bilharzia (schistosomiasis)
  - Ketamine use
  - Ureteric stent

- **Ureteric**
  - Calculus in the lower end
  - Tuberculous ureteritis

- **Prostatic**
  - Acute inflammation
  - Abscess

- **Vesicular**
  - Acute seminal vesiculitis

- **Rectal**
  - Carcinoma

- **Anal**
  - Fissure or ulcer
  - Inflamed haemorrhoids

Diseases of the bladder

A *calculus* in the bladder, unless it is trapped in the pouch behind an enlarged prostate or in a diverticulum, causes pain in the glans penis after micturition. It may exist without causing cystitis, although commonly there is some degree of pyuria. There is increased frequency and urgency of micturition, and often an associated urinary tract infection. The terminal drops of urine during micturition may be blood-stained, and there may have been a sudden reduction in the stream during micturition. In some cases, there is a previous history of acute ureteric colic due to the descent of a stone from the kidney.

The great majority of vesical calculi are radio-opaque and can be seen on a plain X-ray of the pelvis. At cystoscopy, stones can be seen, their approximate size determined, and any other conditions of the bladder accompanying or simulating calculus diagnosed with certainty. Following treatment for the calculus, most patients need treatment for bladder outflow obstruction in the form of a transurethral resection of the prostate or other outflow surgery.

*Bladder tuberculosis* is usually secondary to tuberculous disease in some other part of the genitourinary tract, particularly the kidney. It causes marked penile pain after micturition, together with pyuria and a tinge of blood in the terminal drops of urine; the frequency of micturition is increased during both day and night. In a young patient in whom increased frequency of micturition, pyuria and penile pain are present, a search should be made for any tuberculous focus, especially in the kidneys, by contrast CT or excretion urography, and in the epididymes, prostate or seminal vesicles, or for marked thickening of the terminal ureter as felt per rectum.

Routine laboratory examination of the urine reveals an acid urine, with pus cells but no growth on routine culture – the so-called ‘sterile acid pyuria’. The deposit from three early-morning specimens should be examined for acid-fast bacilli, and polymerase chain reaction or gene probe tests can distinguish *Mycobacterium tuberculosis* from other mycobacteria. A cystoscopic examination with biopsy and culture may be necessary to determine the extent of the disease.

*Bladder tumours*: carcinoma of the bladder occurs in a papillary or a solid form. This begins most commonly in the base of the bladder and can cause trigonal irritation with frequency, urgency and dysuria. The classic symptom is, of course, painless macroscopic haematuria, and all new cases warrant urgent cystoscopy and upper tract imaging. New-onset dysuria with microscopic haematuria in a smoker should always raise the possibility of bladder cancer and its important dangerous precursor, carcinoma-*in-situ*.

Occasionally, bladder cancers may give rise to renal pain when the infiltration has extended to and occluded one or both ureteric orifices. A CT-IVU or the cystogram phase of intravenous urography (IVU) (Figs P.14 and P.15) may sometimes show a filling defect in the otherwise regular contour of the bladder. Cystoscopic examination, together with transurethral resection biopsy, will usually confirm the diagnosis and provide definitive treatment in about 80 per cent of cases.

*Acute cystitis* causes tingling pain in the penis after micturition from the inflammatory infiltration of the trigonal area. The mode of onset, the character of the pain and other symptoms of cystitis will point to the cause of the pain.

*Bilharzia* (schistosomiasis), due to infection with the trematode *Schistosoma haematobium*, gives rise to clinical symptoms very similar to those of vesical tuberculosis. The history of residence in an infected district (e.g. Egypt or East Africa), microscopic examination of the urine for ova, and the typical cystoscopic appearance of the bladder (sandy patches near the trigone) establish the diagnosis. Radiographs
may show calcification of the bladder or ureters, and a CT intravenous pyelogram or standard pyelography often demonstrates stricture formation and gross dilatation of the ureters.

Regular use of the recreational drug ketamine may cause urinary frequency, urgency and post micturition penile pain. This may not be evident from the history unless it is specifically asked about. Irreversible damage can occur leading to a small shrunken bladder.

The distal coil of a ureteric stent in the bladder often causes post-micturition pain relieved only by removal of this foreign body.

**Ureteric lesions**

Ureteric lesions infrequently produce pain in the glans penis after micturition and may cause considerable difficulty in the diagnosis from vesical disease.

When a calculus becomes impacted in the narrowed distal intramural ureter, symptoms are produced almost exactly similar to those of bladder calculus or tuberculosis, namely increased frequency of micturition, referred pain in the glans penis after micturition, and a small amount of pus and blood in the urine. A careful history will often be of value in these cases; the first attack of pain is usually described as being sudden, and felt in the renal angle posteriorly, passing forwards above the iliac crest and spine, and finally becoming localized at the situation of the external abdominal ring. With ureteric calculus, there is usually aching pain in the kidney of the affected side from the dilatation of its pelvis. The gold standard imaging modality is now a non-contrast CT, but a good plain X-ray of the pelvic area as part of an IVU series may show a stone. A total of 90 per cent of ureteric calculi are radio-opaque, but they may be confused with phleboliths or be obscured by gas shadows or underlying bony structures. A cystoscopic examination may be required to exclude a bladder lesion if imaging is not definitive, and marked congestion and dilatation of the blood vessels in the immediate vicinity of the ureteric orifice indicates a stone in the intramural ureter.

Treatment is either conservative with analgesia and alpha-blockers for small calculi, or active by rigid ureteroscopy or extracorporeal shockwave lithotripsy for larger or resistant stones.

In renal tuberculosis, the penile pain and increased frequency of micturition are more marked, the kidney may be felt to be enlarged and tender, and tubercle bacilli will be found in the urine. Apart from this, typical changes in the ureteric orifice are seen on cystoscopic examination, the orifice being pulled up, retracted or

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**Figure P.14** Intravenous urogram taken 20 minutes after injection. There is a filling defect on the right side of the bladder due to a large benign papilliferous tumour. Both kidneys are normal.

**Figure P.15** Intravenous urogram. There is a filling defect on the right side of the bladder caused by infiltrating carcinoma. The right ureteric orifice has been obstructed, and there is no function of the right kidney.
horseshoe-shaped, and usually occupying a position slightly above and outside the situation of the normal orifice, due to the actual shortening of the ureter by infiltration of the submucous coats. The rigid ‘golf-hole’ ureteric orifice is a late manifestation caused by contraction of scar tissue around it and in the ureter above it.

**Diseases of the prostate**

These often cause pain in the penis immediately following micturition. This is seen most commonly with acute inflammation or abscess in the gland as a sequela of acute gonorrhoea or septic urethritis. In either case, there is penile pain, sometimes associated with erection, but little difficulty will be experienced in the diagnosis due to the specific symptoms and after rectal examination.

**Diseases of the seminal vesicles**

These are seldom present without accompanying disease of the prostate or bladder. Acute vesiculitis may follow urethritis and give rise to pain after micturition, but in most cases it will be associated with prostatitis. Similarly, tuberculous nodules in the vesicle will be associated with foci in the epididymis, prostate or bladder.

**Diseases of the rectum and anus**

These may occasionally give rise to penile pain following micturition, apart from any infection of the bladder or prostate. Thus, an infiltrating carcinoma in the anal canal, an anal fissure or an inflamed haemorrhoid may occasionally cause pain in the penis, but in each the local symptoms of the trouble will be the more marked, and little difficulty will be found in the diagnosis if a local examination is made with care.

**PAIN IN THE PENIS APART FROM MICTURITION**

These include certain lesions of the penis and urethra, and also the pains referred from disease elsewhere. Although a local lesion may cause little more than discomfort in many patients, in some it is described as pain, the degree of which depends upon the nervous susceptibility of the individual. Thus, penile pain may be present with acute urethritis, with balanitis in association with phimosis or with paraphimosis. In some instances, herpes of the prepuce or penile skin causes distinct pain. Any infiltration of the cavernous tissue of the penis causes pain during erection of the organ; therefore, during an attack of acute urethritis, the symptom known as chordee arises from this cause. It may occur in a chronic form in Peyronie's disease, a condition of unknown aetiology but sometimes associated with Dupuytren's contracture and retroperitoneal fibrosis. In this condition, erection is not only painful but may be accompanied by lateral deviation of the organ due to fibrous plaques in the corpora cavernosa. Another condition causing the same trouble arises from haematoma in the cavernous tissues of the penis following a local injury, due either to external violence or arising during forcible attempts at coitus. A similar condition may arise spontaneously in blood diseases, especially lymphatic or myelocytic leukaemia. Squamous cell carcinoma of the penis on rare occasions gives rise to pain in the organ.

Finally, pain in the penis may be based on an anxiety state or some other mental cause rather than organic disease.

**PERINEUM, PAIN IN**

Harold Ellis

Pain in the perineum is a symptom often mentioned by patients in giving their history of some affection of the genitourinary apparatus or of other organs, but usually only as a dull aching, of which little notice is taken, as it is generally of minor degree in comparison with other more striking symptoms. The complaint of perineal pain per se does not convey much information to the clinician, and it is practically never present as the only symptom in a case. It may be a manifestation of an anxiety state.

Aching in the perineum is frequently present in diseases of the following organs:

- **Prostate**
  - Chronic prostatitis
  - Abscess
  - Calculus
  - Adenomatous enlargement
  - Carcinoma
- **Seminal vesicles**
  - Acute inflammation
  - Tuberculosis
- **Testis**
  - Congenital misplacement in the perineum
- **Urinary bladder**
  - Cystitis
  - Tuberculosis
  - Calculus
  - Carcinoma
- **Urethra**
  - Injury
  - Gonorrhoea
  - Stricture with extravasation or urethral abscess
  - Fistula
  - Calculus impacted in the bulbo-prostatic portion
• Anal area
  – Haemorrhoids
  – Fissure
  – Carcinoma
• Vagina
  – Acute inflammation
  – Inflammation or abscess of the Bartholin’s glands
  – Cystocele
  – Epithelioma
• Cutaneous diseases
  – Intertrigo
  – Diabetic inflammation
  – Condylomas

From the foregoing list, it will be seen that aching in the perineum occurs with numerous different lesions, but other symptoms discussed elsewhere are in almost every case more marked. In prostatic disease, it is an indication of inflammation rather than of enlargement. In clinical practice, it is most commonly found to be due to chronic prostatitis. Examination of the secretion expressed after prostatic massage will show the presence of many pus cells.

PERINEUM, ULCERATION IN
Harold Ellis

Ulceration may be present in the perineum as the result of:
• Cutaneous inflammation or injury
• Urethral suppurations or fistulas
• Prostatic supputation
• Anal fistula
• Syphilis
• Granuloma venereum (inguinale)
• Lymphogranuloma inguinale (venereum)
• Epithelioma and other cutaneous cancers

CUTANEOUS INFLAMMATION OR INJURY
An ulcer in the perineum may result from direct injury to the area, or from infection of the sebaceous or hair follicles. An ulcer from these causes may be placed at the centre or to one side of the perineum, is movable on the deeper parts, and shows no track into which a probe can be passed. In women, ulceration of the perineal area may be associated with gonorrhoeal or septic vaginal discharge. It may also arise from severe scratching caused by the irritation of such skin infections as tinea cruris or pruritus ani.

URETHRAL SUPPURATIONS OR FISTULAS
During the progress of an acute urethritis, a glandular follicle may become infected. The suppurative process leading from this in the bulbous urethra may extend towards the perineum and open externally, leaving a small fistula that may or may not discharge urine during the act of micturition. In a similar manner, urinary fistulas may result from inflammatory processes behind a urethral stricture, and in an old-standing case it is not uncommon to find a urinary calculus in the dilated portion of the urethra behind the stricture. Where the urethral suppuration is acute and an abscess bursts in the perineum, the diagnosis will be obvious, and the ordinary treatment for an abscess, in addition to that of the acute urethritis, will usually suffice to cure the condition. If the perineal wound discharges urine, this occurs usually only during the act of micturition, as there is no interference with the vesical sphincter. A stricture of the urethra, not necessarily of sufficient degree to cause severe interference with micturition, will generally be seen on endoscopic examination, the sloughy granulations behind it denoting the position of the urethral opening of the fistula.

DISEASES OF THE PROSTATE
An abscess or tuberculous focus in the prostate may occasionally discharge in the perineum and remain as a sinus. An abscess in the prostate practically always arises from some infection in the posterior urethra, from venereal causes or after septic instrumentation. It is accompanied by urethral discharge, or there is a history of a recent infection, while per rectum the prostate may be felt to be inflamed, or scarred from the shrinkage of the abscess cavity.

ANAL FISTULA
An ulcer on the perineum may be present as the result of an anal fistula – commonly from perianal supputation and occasionally as a tuberculous infection. The history of pain on defecation followed by the rupture of an abscess and the history of passage of flatus or faecal matter from the fistula are usually present, or a probe may be passed into the fistula and felt by a finger passed into the rectum. Perianal and perirectal abscesses, fissures and fistulas may occur in Crohn’s disease, especially when the colon is involved, and less commonly in ulcerative colitis (see Fig. R.3).

SYPHILIS
Syphilis may cause ulceration on the perineum either as a chancre or as mucous tubercles. A chancre at this site is rare. It forms a small ulcer with slightly indurated borders, indolent in character, and accompanied by slight enlargement of the inguinal lymph nodes. A chancre of the skin may not possess the usual features of a genital chancre, and it is not usually diagnosed with certainty until the secondary
lesions of syphilis become apparent; but an ulcer with raised, infiltrated edges, which shows no tendency to heal under aseptic precautions, should always give rise to a suspicion of syphilis. Treponema pallidum should be looked for, under dark-ground illumination, and serological tests for syphilis should be performed.

Condyloma lata may be present about the perineum in association with active syphilis. They may extend from the anal or vulval orifice, and form oval or rounded, flat-topped, sessile masses, covered by macerated greyish epithelium, or they may be ulcerated on the surface. The accompanying signs of syphilis will indicate the diagnosis.

Soft sores may occur in the perineum as well as on the scrotum or the vulva; they are generally venereal but are not in themselves syphilitic; they are generally multiple, are apt to be foul, and cultures from them yield Ducrey's bacillus (Haemophilus ducreyi).

ULCERATION OF GRANULOMA INGUINALE

Ulceration of granuloma inguinale sometimes attacks the perineum, and fistulas there can be caused by lymphogranuloma venereum (see PENILE LESIONS, p. 502; see also Fig. R.2).

EPITHELIOMATOUS ULCERATION

Epitheliomatous ulceration of the perineum is seen as a direct spread of a growth of the anus or vulval area, when the diagnosis presents no difficulty. An epithelioma may develop in the scar of some former cutaneous affection, particularly in long-standing fistula-in-ano, in which case an ulceration may exist showing the usual characteristics of a cutaneous epithelioma. The inguinal nodes may be enlarged early from the inflammatory process, or later by invasion with malignant disease. Other cutaneous cancers, malignant melanoma and basal cell carcinoma may also occur in this situation. In case of doubt, a biopsy specimen should be taken for microscopical examination.

PERISTALSIS, VISIBLE

Harold Ellis

(See also BORBORYGMI, p. 69.)

Visible peristalsis is usually pathological. However, in a number of conditions, the normal movements of the bowel may be visible; these circumstances are divarication of the abdominal recti muscles, an incisional or massive umbilical hernia containing bowel, and extreme thinness of the abdominal parietes – the result of emaciation or, rarely, congenital absence of the recti. It is not uncommon to see visible peristalsis within the sac of a very large ventral or inguinoscrotal hernia. In all these circumstances, the diagnosis can be made at inspection, and the patient is otherwise symptomless. In all other situations, visible peristalsis is pathological and may be of two types, gastric and intestinal.

GASTRIC PERISTALSIS

Gastric peristalsis takes the form of a comparatively large swelling in the upper abdomen showing slow waves of peristalsis that progress from under the region of the left ribs, slowing downwards and to the right. This swelling indicates obstruction to the gastric outlet. There may be other signs of gastric dilatation and distension, particularly a loud succession splash (Fig. P.17). Typically, there is a history of the vomiting of large amounts of liquid in a projectile manner, which may contain fragments of food ingested 24 hours or more previously. The diagnosis can be confirmed by the passage of a nasogastric tube, which will yield a pint or more of fluid several hours after the last food or drink has been taken. The aspirate has a typical stale, unpleasant smell, and it may contain recognizable particles of food eaten even several days before. A barium X-ray examination will clinch the diagnosis by demonstrating the gastric retention and dilatation. An X-ray taken 6–8 hours after the ingestion of the barium is particularly valuable as this will confirm the extent of gastric hold-up (Fig. P.18). In doubtful cases of visible gastric peristalsis, the sign may be accentuated by asking the patient to swallow several glasses of soda water: in the normal subject, no peristalsis is seen; in cases of pyloric obstruction, previously invisible peristalsis may now become obvious.
In congenital hypertrophic pyloric stenosis of infancy, not only can gastric peristalsis be seen after a drink from a bottle, but the hypertrophied pylorus can often also be felt. This interesting and eminently treatable condition does not become apparent immediately, but around 4 weeks after birth.

**VISIBLE INTESTINAL PERISTALSIS**

Visible intestinal peristalsis is a feature of advanced intestinal obstruction, with the limitations discussed above. As a pathological entity, it will not occur alone but is accompanied by colicky abdominal pain, abdominal distension, vomiting and absolute constipation. The discussion of the differential diagnosis of the different causes of the symptoms will be found elsewhere. If the small intestine alone is involved, the waves are multiple and run more or less transversely across the abdomen – the ladder pattern. When the colon is obstructed, peristalsis takes the form of vertical waves, especially in one or both flanks, but this is much more rarely seen.

Plain radiographs of the abdomen taken in the erect and supine positions are invaluable; the first demonstrates multiple fluid levels, the second the distribution of gas shadows within the dilated loops of bowel, which will often enable the clinician to determine whether small or large bowel is obstructed.

**PHOBIAS**

Andrew Hodgkiss

A phobia is a persistent, irrational, morbid fear of a specific object, situation or activity that induces a compelling desire to avoid that stimulus. This reaction is acknowledged as inappropriate or disproportionate, but nevertheless the individual is unable to desist from avoidance behaviour because of the anxiety that develops when exposure occurs or is anticipated. The causes of phobias are listed in Box P.3.

Most people probably harbour mild phobic responses to such common stimuli as insects and snakes, spiders, dogs, small furry animals, lifts, heights, air travel, darkness, tunnels, blood, vomiting, dentists – and even doctors! All are catalogued by classically derived prefixes that can be employed to impress. Typically, these exaggerated fears can be overcome if necessary, place no limitations upon lifestyle and are often regarded as socially acceptable reactions, so treatment is rarely sought or required. Clinically appreciable phobias are reported in about 8 per cent of the adult population, about 1 in 40 cases being severely disabling.
is distinguishable from obsessional ruminations often signify an underlying stress. The presentation hypochondriacal traits in their personality, and they common in patients with appreciable obsessional or radiation and food additives. Illness phobias are more attention, hence topical illness phobias including AIDS, implicated – and particularly when attracting media venereal disease, but any disorder can become illness phobias concern cancer, heart disease and is already convinced the disease is present. Common not have the disease, which contrasts importantly will seek reassurance from the doctor that they do from a disease rather than primary avoidance. Patients endless ruminations about the possibility of suffering Phobias differ in that the phobia consists of thoughts, and from hypochondriasis in that there is a single concern with reassurance often effective. Illness phobias can be the presenting feature of a primary depressive illness when they respond to antidepressant treatment but, if unrecognized, illness phobias may progress to frank hypochondriacal delusions in depressed patients.

AGORAPHOBIA

While simple phobias are the most common form of the disorder, agoraphobia tends to be the most incapacitating, and it accounts for over half the phobic presentations to specialists. The essential features of agoraphobia are a marked fear of being alone in public, accompanied by apprehension of becoming helpless in, or unable to escape from, a crowded public place. Life becomes increasingly constricted as fears about streets, travelling, shopping and crowded spaces take root. Typically, the patient will battle unsuccessfully, and present when virtually restricted to their home or dependent upon others for accomplishing routine daily activities. This condition is twice as common in women as men, usually develops in early adult life, and is associated with personality traits of passivity, dependence and anxiousness. A precipitant that may threaten the individual’s security is often evident – commonly a social change such as marriage, divorce, bereavement or childbirth; less commonly an incapacitating, unpredictably episodic illness such as asthma, epilepsy or Ménière’s disease; and sometimes a more persistent physical change, for example following a head injury or other brain disease, amputation, transplantation, stoma formation or other form of major surgery. Once established for about a year, agoraphobia usually persists with remissions and relapses linked to life stresses.

Agoraphobia is almost always accompanied by other neurotic features – panic attacks, free-floating anxiety, depression, hyperventilation, obsessions and depersonalization are all frequently reported. As with simple phobias, schizophrenia or obsessive–compulsive disorder may occasionally be underlying, but the most common cause of secondary agoraphobia is depressive illness.

Sometimes it can be impossible to decide clinically whether the patient has depression with secondary agoraphobia or agoraphobia with secondary depression, although evidence of agoraphobic changes over months or years would point to the latter. In circumstances of uncertainty, it is worthwhile prescribing an antidepressant drug as secondary agoraphobia will remit.
Alcohol dependence syndrome is another condition that can present with agoraphobia, so a careful alcohol history is an important part of the assessment.

Finally, it is advisable to inquire about health and education problems in the children of agoraphobic women. Children may not be going to school either because their mother would be stranded at school or, more seriously, requires them as a crutch at home; sometimes school phobia or other emotional problems emerge, and occasionally the child can form the initial presentation.

SOCIAL PHOBIAS

Social phobias are especially common in adolescents. The patient avoids social situations because of the intense fear of behaving in an embarrassing or humiliating manner, and the feeling of being under scrutiny by an audience who are able to detect minor signs of their anxiety. Common social phobias are speaking, eating, drinking, writing or blushing in public, and using public lavatories. Usually, the patient has a single phobia linked to a specific situation, but occasionally all forms of social contact are avoided, and the patient leads the life of a recluse.

Many social phobics treat their fear and anxiety by using alcohol or tranquillizers, and consequently some present with alcohol or drug dependence. Other phobias can co-exist, although agoraphobia tends to develop later in life: incidentally, agoraphobics also fear groups of people, but their phobia centres upon the mass of people, while social phobics are apprehensive of the individuals within the crowd. Again, schizophrenia and obsessive–compulsive disorder may underlie a social phobia, although in depressive illness unresisted social withdrawal is more common and is motivated by anhedonia rather than fear. In more profound social phobia, the possibility of a developing schizophrenic illness should be kept in mind, particularly if the patient is less concerned about their impoverished lifestyle than would be anticipated. For some patients with avoidant personality disorder, social phobias establish a persistent, unwelcome impairment of life, with work and relationships grossly affected. However, such patients are rare, and the majority of social phobics have an excellent outlook, and they often with a minimum of advice and assistance.

PHOTOPHOBIA

David Werring & Mark Kinirons

Photophobia is an intolerance of light. The causes can be grouped into ocular and non-ocular. Ocular causes may be due to lack of retinal pigment, ocular inflammation or other ocular damage. Reduced retinal pigment occurs in cone dystrophies and oculocutaneous albinism. Chediak–Higashi syndrome is a rare autosomal recessive disorder characterized by recurrent pyogenic infections, partial oculocutaneous albinism, progressive neurologic abnormalities and mild coagulation defects. Causes of ocular inflammation include uveitis and keratitis. Some patients complain of photophobia in an eye with a mydriatic pupil because of the increased light through the wider aperture. Other ocular causes of photophobia include corneal abrasion, congestive glaucoma, retinal detachment and refractive surgery. The diagnosis can be made by detailed ophthalmological assessment.

Migraine is commonly associated with photophobia and can usually be diagnosed by the accompanying features; these include lateralized, throbbing headache with phonophobia (intolerance of sounds), a desire to lie still, nausea and sometimes vomiting. Photophobia is an important component of the syndrome of meningeal irritation, together with neck stiffness and positive Kernig’s and Brudzinski’s signs. Kernig’s sign is limitation or pain on extending the knee with the leg flexed at the hip; Brudzinski’s sign is flexion of the hips when the head is pushed forward on to the chest. Meningeal irritation may be due to meningitis (acute or chronic), encephalitis or subarachnoid haemorrhage. The diagnosis should be suggested by the associated neurological symptoms and signs as well as investigations including brain imaging and lumbar puncture.

PICA

Dipak Kanabar

Pica is a persistent pattern of eating non-nutritive substances (e.g. dirt, paper, plaster, chalk), lasting 1 month or longer. It is more commonly in young children with 10–32 per cent of children aged 1–6 years old exhibiting these behaviours. Clay, dirt, ice, sand, animal faeces, paint and hair balls are just a few examples of what children and adults with pica have been known to eat.

Pica is seen in children with emotional deprivation, neglect or even child abuse, especially between the ages of 2 and 5 years; it is interpreted as the child’s symbolic response to a lack of maternal affection and care. It may lead to poisoning if noxious substances are ingested accidentally (e.g. lead), or to obstruction of the stomach if multiple foreign bodies are eaten (e.g. pebbles). If the swallowed material is fibrous, it may bind together to form a bezoar, which forms a cast of the stomach with consequent obstruction. A variant...
of pica is *trichotillomania*, where hair is pulled out and eaten with the formation of a trichobezoar.

There is no single test that confirms pica. However, since pica is associated with abnormal nutrient or raised toxic metal levels, and in some cases malnutrition, several tests may be performed. Serum levels of lead, iron and zinc should be taken. Haemoglobin should also be checked to test for anaemia. Lead levels should always be checked in children, who may have eaten paint or objects covered in lead paint dust. The presence of infection may be detected if contaminated soil or animal waste is being ingested.

Identified nutritional deficiencies and other problems, such as lead toxicity, should be addressed medically. Treatment emphasizes psychosocial, environmental and family guidance approaches. Other successful treatments have been mild aversion therapy followed by positive reinforcement. Medication may be helpful in reducing the abnormal eating behaviour if pica occurs in the course of a developmental disorder, such as mental retardation, or pervasive developmental disorder.

**PILIMICTION**

Harold Ellis

Pilimiction – i.e. the passage of hairs in the urine – is a rare condition that almost invariably signifies that the patient has a pelvic dermoid cyst that has become inflamed, thereafter opening into the bladder and discharging its contents via the urinary passages. This condition has been observed in men, but it occurs more often in women. Subacute or acute cystitis accompanies the event with vesical pain, frequency of micturition and pyuria. The obvious fallacy in diagnosis arises from the possibility of contamination in the urine of hairs that were not, as supposed, passed per urethram.

**PLEURAL EFFUSION**

Alex West

Pleural effusion – the presence of fluid lying between the visceral and parietal pleura – is common and may be associated with a large number of conditions. Of these, congestive cardiac failure is the commonest, and malignant pleural infiltration and infection are the next two commonest.

**PHYSIOLOGY OF THE PLEURA**

In health, the two pleural surfaces are in close contact but separated by a thin layer of fluid. Estimates of the volume vary, but quoted figures range from 1 to 20 ml, with an electrolyte content similar to that of serum and a low protein concentration. The fluid is formed by transudation from the parietal pleura and is absorbed by the visceral pleura. It is in a dynamic state, with about two-thirds of the fluid being absorbed and replaced every hour.

The pleura transmits the forces generated by the respiratory muscles to the lungs, and there is a negative pressure within the pleural space of about −5 mmHg. Capillary fluid and gas would enter the pleural space were it not for a number of balancing factors, including a hydrostatic pressure difference between the parietal capillaries and the capillaries of the visceral pleura, which are supplied by the low-pressure pulmonary arterial system. Plasma oncotic pressure is the same in both sets of capillaries (about 35 mmHg), while pleural osmotic pressure is only about 6 mmHg due to its low protein content. Thus, fluid is driven in sequence from the parietal pleural capillaries to the pleural space, and then on to the visceral pleural capillaries and lymphatics, resulting in a continuous transfer of low-protein fluid.

In the case of gas entering the pleural space, there is a driving force of about 40 mmHg (atmospheric pressure − pleural pressure + pleural capillary blood gas tension) that assists gas absorption, as occurs in closed pneumothoraces.

**CLINICAL FEATURES**

The symptoms associated with the accumulation of fluid in the pleural space depend upon the cause, volume and rate of formation of fluid. Small effusions are often symptomless, and even quite large effusions can cause little disability, provided the fluid has accumulated slowly. Effusions caused by inflammatory disease often present with pleuritic pain, which may be relieved as the fluid accumulates. Large effusions eventually cause symptoms including shortness of breath, initially on exercise and later at rest, together with dull, aching discomfort or pressure over the affected side of the chest.

The clinical findings are influenced by the size and site of the effusion. Most effusions occupy the dependent part of the pleural space, so when the patient is sitting, the characteristic findings of stony dullness to percussion and distant or absent breath sounds are most prominent at the lung bases. Bronchial breath sounds or aegophony may be heard directly above an effusion. Large effusions displace the mediastinum towards the unaffected side unless the underlying lung is fibrosed from previous inflammation (tuberculosis) or collapsed due to a proximal bronchial lesion. Very large effusions may displace the mediastinal contents...
to produce an area of dullness at the opposite base close to the midline (Grocco’s sign).

RADIOLOGICAL FEATURES
Effusions may be small, moderate, large, encysted, mediastinal or subpleural. Small effusions may be difficult to detect clinically but can be seen radiographically (Fig. P.19) as non-specific blunting of the costophrenic angles. Moderate-sized free effusions cast a characteristic homogeneous shadow over the lower lung fields, obscuring the diaphragm and cardiac silhouette. At the upper border of the effusion, there is decreased density of the radiographic opacification, with a superior concave curvature that appears to reach its highest level in the axilla when seen on posteroanterior films, or posteriorly on lateral films. In fact, the level of the effusion is horizontal, and the radiographic appearances are artefactual due to the increased distance traversed by the radiation. Massive effusions cause complete opacification of the hemithorax, often with very marked displacement of the mediastinal structures to the opposite side. Sometimes the fluid collects in the pleural cavity under the lung adjacent to the diaphragm, the so-called diaphragmatic or subpulmonary pleural effusion. The upper margin of the fluid shadow runs parallel to, and may be mistaken for, an elevated diaphragm. On the left side, the apparent separation of the transradiant gastric ‘air bubble’ from the transradiant lung tissue may draw attention to the effusion. If the presence of fluid is in doubt, a lateral decubitus film may help to differentiate between effusion or pleural thickening when, in the former, the fluid is seen to shift to the lateral chest wall or mediastinum. Interlobar pleural effusions are quite commonly seen as an extension of an effusion, and they result in a characteristic ovoid homogeneous shadowing with well-demarcated margins lying in one of the interlobar fissures. Interlobar effusions may mimic tumours and occur particularly in cardiac failure when clearance following diuretic treatment gives rise to the term ‘vanishing pulmonary tumour’.

Encysted effusions can give rise to diagnostic difficulties, especially if they lie posteriorly and cause homogeneous shadowing suggesting consolidation when seen on posteroanterior films. The nature of the abnormality becomes apparent in a lateral view when opacification is seen to lie posteriorly. Although the chest X-ray gives the diagnosis, it can be misleading. Pleural ultrasound and/or CT scanning are far more accurate and will add more information on the effusion, including its characteristics, and they will often help establish the underlying cause.

AETIOLOGY
Pleural effusions can be divided into transudates and exudates. The pleural fluid formed through normal capillary membranes is a transudate with a low protein content, whereas fluid formed through abnormally permeable capillary walls contains a higher concentration of protein. The distinction between a transudate and exudate may be obvious on clinical grounds, for instance when there is cardiac, renal or hepatic failure. To distinguish between the two, ‘Light’s criteria’ should be used on the sampled fluid, assuming that the macroscopic appearance does not confirm another diagnosis (see below). This involves measuring the pleural fluid total protein and lactate dehydrogenase (LDH) level. There are three criteria and, if any one of these is positive, the fluid should be considered an exudate and thus may require further investigation.

1. If the ratio of pleural fluid total protein to serum is >0.5
2. If the ratio of pleural fluid LDH to serum is >0.6
3. If the pleural fluid LDH level is greater than two-thirds of the upper limit of normal of the hospital’s serum LDH reference range

The majority of exudative pleural effusions are related to increased pleural capillary permeability and occur in response to inflammation (both infective and other), ischaemia (pulmonary infarction) or pleural neoplasia.

Figure P.19 Right pleural effusion (white arrow) and pulmonary plethora (black arrows) from heart failure in a fluid-overloaded renal dialysis patient.
The clinical findings may provide some diagnostic clues, as may radiology, which will also localize the position of the fluid and reveal its extent. In most cases, the fluid must be examined cytologically and biochemically, and be cultured for organisms. The appearance of the freshly aspirated fluid can be informative. It is usually straw-coloured or cloudy, but it can be frankly purulent (empyema), blood-stained (haemothorax) or opalescent (chylolithorax). Blood-stained effusions are relatively common and, setting aside trauma to the chest or accidental haemorrhage resulting from a traumatic pleural tap, are most commonly seen in malignancy or pulmonary infarction. A haemacrit on heavily blood-stained fluid can be compared to peripheral blood and, if similar, may confirm trauma as the cause.

The pleural fluid cytological findings in benign effusions are variable. Mesothelial cells and macrophages are found in most transudates. Mesothelial cells predominate in exudates, especially when due to pulmonary infarction. Neutrophil polymorphonuclear leucocytes are frequently present in sterile and infected effusions associated with pulmonary inflammation. The exception to this pattern is tuberculosis, where an initial neutrophil excess is quickly replaced by lymphocytes, which may account for between 80 and 100 per cent of all the cells present in the effusion. Lymphocytic pleural effusions are not specific to tuberculosis and also occur in some forms of malignancy (e.g. lymphoma) or in chronic effusions of heart failure.

A high pleural fluid eosinophil count is seldom associated with allergy, but it is seen in occasional pulmonary infarcts or malignancy, or simply when there has been bleeding, instrumentation or air allowed into the pleural space.

Malignant cells are present in about 30–60 per cent of cases where there is malignant invasion of the pleura. The appearances are often diagnostic of the cell type and, when due to metastases, often can reveal the site from where the primary tumour has arisen. Other useful characteristics of the fluid include measurements of pH and blood glucose. A low pH (pH less than 7.20) is occasionally seen in tuberculosis but is more helpful in parapneumonic effusions prior to the onset of frank empyema where this is an indication for formal drainage. The presence of a low pleural fluid glucose level may well be seen in the presence of infection. A very low fluid glucose suggests rheumatoid pleural disease in the right clinical context, although rarer cases of tuberculosis and malignancy may also be found to have a low pleural fluid glucose.

The diagnostic sign of pleurisy is a rub of a creaking superficial nature, usually located close to the site of pain. The pleura is a double serous membrane separating the lung from the chest wall and mediastinum. The pleural surface consists of a uniform layer of mesothelial cells supported on a connective tissue framework well supplied with capillary and lymph vessels. The parietal pleura is innervated with pain-sensitive nerve fibres supplied by the intercostal and phrenic nerves, and it is exquisitely sensitive to painful stimuli. In health, the visceral and parietal pleural surfaces are smooth, glistening and separated by a small amount of fluid, allowing low friction movement of the lungs with respiration.

In contrast, if the pleural surfaces become thickened or roughened by inflammation or neoplastic infiltration, movement with breathing will cause increased friction, which may be heard with a stethoscope. Patients commonly complain of thoracic pain on breathing or coughing. On auscultation, sounds of varying intensity may be heard during inspiration and expiration, often described as having a leathery or creaking quality that may be exaggerated when the stethoscope is applied firmly to the chest wall. Pain associated with a pleural rub may vary in degree from lancinating discomfort during slight inspiratory effort to a less sharp ‘catch’ of pain at the end of maximum inspiration. Pleural pain is often reduced by breath-holding or exerting firm pressure over the affected thoracic segment. Except when it involves the diaphragm, the affected pleura typically underlies the area in which pain is perceived. The central portion of the diaphragm is innervated from the third and fourth cervical posterior nerve roots running via the phrenic nerve. Pain caused by diaphragmatic pleural irritation is often referred to the neck and shoulders.

A pleural rub may last for as little as a few hours in short-lived inflammatory conditions, such as pneumonia, to months or even years in patients with more chronic causes of pleurisy. Typically, as the pleurisy settles, the pain and physical signs, including the rub, become less obvious, although in some patients the rub may persist after the pain has gone, and occasionally a loud rub may persist indefinitely.

Most pleuritic conditions giving rise to pain and an auscultatory rub are inflammatory in origin. Infection associated with community-acquired pneumonias, especially pneumococcal, *Mycoplasma* and other
‘atypical’ infections, may present with severe pleurisy and a pleural rub accompanied by signs of pneumatic consolidation. Pulmonary infarction secondary to pulmonary embolism is another frequent cause, and this is especially common following major surgery, in patients with underlying malignancy, when thromboembolism may be the first sign, or in females taking oestrogen preparations. Tumours invading the chest wall typically cause a continuous persistent pain but may occasionally present with pleurisy and a pleural rub. Rather less frequently, a pleural rub may occur in association with asbestos-induced pleural disease, or connective tissue diseases such as systemic lupus erythematosus or rheumatoid arthritis. Recurrent pleurisy at the same site should suggest bronchiectasis, and that at different sites, bronchopulmonary aspergillosis. If pleurisy progresses to a pleural effusion, the sharp pain and pleural rub largely disappear, to be replaced by a dull and more constant ache and heaviness.

The pain of pleurisy may be mimicked by a number of chest wall conditions, such as rib fractures, intercostal muscle pain due to tearing or strain, Tietze’s syndrome, and neurogenic causes such as intercostal nerve root compression and herpes zoster. Pain due to intercostal muscle strain and tears can be quite sharp, may be caused by coughing, and can result in shallow breathing. However, local tenderness over the affected site is common, and typically no pleural rub is heard.

Epidemic myalgia (pleurodynia, Bornholm disease, devil’s grip or epidemic dry pleurisy) is an acute febrile viral illness affecting skeletal muscle characterized by an abrupt onset of intense pain in the lower chest or upper abdomen. In about 25 per cent of patients, headache, malaise, anorexia, sore throat and deep myalgia precede the onset by 1 or 2 days. The afflicted patient complains of a fever of 38–40 °C and multiple paroxysms of excruciating pain lasting from a few minutes to several hours. The illness is often biphasic, with an initial bout of pain and fever settling, only to recur after a day or two. The acute illness usually settles within a week, but rarely patients have several recurrences over a period of several weeks. The illness may be accompanied by myocarditis or pericarditis.

Epidemic myalgia is caused by enteroviruses, usually Coxackie B3 or B5, but also Coxackie A or echoviruses. The incubation period is short, about 3–5 days, and, as with other entroviral infections, the majority of illnesses occur in the summer and autumn. A specific diagnosis can be made by isolating virus from the throat and faeces during the acute illness, or demonstrating a rising titre of serotype-specific neutralizing antibodies in acute and convalescent sera. The level of creatine phosphokinase in the serum may also be elevated, reflecting injury to striated muscle. Confusion with acute myocardial infarction is inevitable in those patients presenting with abnormal electrocardiograms and raised creatine phosphokinase. The condition may also be confused with pre-eruptive herpes zoster although, in the latter condition, pain is more constant and no pleural rub is detected.

Recurrent polyserositis (familial Mediterranean fever) is an autosomal recessive, recurrent inflammatory disease of unknown cause, characterized by recurrent inflammation of the serous membranes. Attacks occur at irregular intervals from several days to several years, with pleurisy, abdominal and joint pain, and other systemic symptoms that typically settle spontaneously within 12–48 hours. This condition usually manifests in children, is recognized in many parts of the world, but is largely restricted to ethnic groups originating in the eastern Mediterranean area.

Investigation of patients presenting with pleurisy and a pleural rub will almost inevitably include a chest radiograph, which is frequently useful in showing a primary lung condition. If the chest X-ray is normal, or if it shows only a small pleural reaction, it may be important to consider the possibility of a pulmonary embolism, and further examination of the legs, together with scanning, may help in coming to a therapeutic decision. If pulmonary embolism is considered unlikely after clinical assessment and testing, and in the absence of any other features, it is reasonable to make a provisional diagnosis of viral pleurisy, and to treat the patient with adequate analgesia.

PNEUMATURIA

Ben Challacombe

The passage of gas per urethram, with, after or independently of urination, is a rare but striking symptom, particularly when it occurs in males. Patients describe passing wind or distinct bubbles urethrally. It should always be taken seriously and appropriately investigated as it is likely to be related to significant pelvic pathology. It may be due to one or other of two distinct groups of causes:

- A fistulous communication between the rectum, caecum, small bowel, appendix or other part of the alimentary canal and the bladder, ureter or renal pelvis, either directly or via an intermediate gas-containing abscess cavity
- More rarely, infection of the bladder or other part of the urinary tract by gas-producing organisms
POLYURIA

In the first group, the patient is likely to pass faecal material as well as gas and has often been recently treated for a urinary tract infection. Lack of faecal material in the urine does not exclude a fistula between some part of the alimentary canal and the urinary tract; the fistula may be small, permitting gas but not faeces to traverse it. Appendicitis or sigmoid diverticulitis may lead to the formation of a local gas-containing abscess containing Escherichia coli or Aspergillus that may open into the bladder and cause the discharge of pus and gas per urethram.

Common pathologies responsible for a recto- or enterovesical fistula are:
- Colorectal adenocarcinoma
- Diverticular disease
- Inflammatory bowel disease, commonly Crohn’s
- Post pelvic radiotherapy, cryotherapy or high-intensity focused ultrasound (HIFU)
- Urological malignancy: bladder, rarely prostate, renal pelvis or ureter
- Pelvic trauma: penetrating or due to bone fragments
- Iatrogenic trauma following abdominal/pelvic surgery
- Appendicitis, with or without associated abscess

Urine should be sent for microscopy and culture. Investigation, after appropriate antibiotic treatment of the infection, involves flexible or rigid cystoscopy plus biopsy, with a fistulous track commonly identified in the left lower wall of the bladder, or an area of acute cystitis seen in this area. In addition, a contrast-enhanced computed tomography (CT) intravenous pyelogram/CT of the abdomen and pelvis and/or colonoscopy/barium enema will be required. If no sign of a fistulous communication between any part of the bowel or a gas-containing abscess cavity with the urinary tract can be identified, the pneumaturia may be considered to be solely due to infection. Such patients are usually elderly women with diabetes who are infected with Escherichia coli, Aspergillus aerogenes, yeasts or a combination of these.

POLYDIPSIA

Polydipsia is defined as excessive or abnormal thirst with consequent increase in fluid intake. The primary stimulus to the sensation of thirst is dehydration, which gives rise to an increase in the plasma osmolality of the blood passing through the thirst centre in the hypothalamus. An increase in plasma osmolality can also be achieved by increasing the solute load, for example by drinking salt water. The sensation of thirst must be distinguished from a dry mouth caused by Sjögren’s syndrome, or mouth-breathing or by drugs (in particular antidepressant and antipsychotic medications with pronounced antimuscarinic activity). Apparent thirst may also be due to a psychiatric disorder, when it is called psychogenic polydipsia.

True polydipsia due to dehydration may be associated with disorders that cause polyuria (see below), such as diabetes mellitus, cranial diabetes insipidus, nephrogenic diabetes insipidus and diuretic therapy. Other causes of dehydration not associated with polyuria include inadequate fluid intake, excessive loss of fluid from the skin (fever, thyrotoxicosis or burns injuries), from the stomach (repeated vomiting), from the bowel (diarrhoea) and into serous membrane-lined cavities, as in acute peritonitis.

POLYURIA

Paul Carroll

The term ‘polyuria’ signifies a larger than normal daily volume of urine. There is considerable variation from subject to subject in the amount of urine passed, but a urinary output of more than 3 litres per 24 hours is nearly always abnormal. Polyuria must not be confused with frequency of micturition due, for example, to prostatic hypertrophy or cystitis. Also, although polyuria will almost always lead to a complaint of nocturia, many individuals complaining of the latter have no increase in the total output of urine but show a reversal of the normal diurnal variation in urine flow. This is the case in most cases of sodium and water retention, as in cardiac failure or the nephrotic syndrome, in adrenal gland disorders and in chronic renal failure. Polyuria may be due either to an increased solute load with obligatory water loss or to a primary water diuresis, and will be discussed under these headings. The main causes are summarized in Box P4.

**Box P.4 Causes of polyuria**

<table>
<thead>
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Other causes

- Following fevers
- After attacks of migraine, etc.
- Paroxysmal tachycardia

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POLYURIA DUE TO INCREASED SOLUTE LOAD
Any osmotically active solute will produce a diuresis if present in excess in the distal tubular fluid. For example, the massive protein breakdown occurring in a large haematoma may be associated with a diuresis, urea itself being the active solute. It is also the mechanism of diuretic therapy in which the solute concerned is sodium. Diabetes mellitus is much the most common pathological condition in which this type of polyuria occurs. The daily urine volume is often 4 litres or more, and polyuria and excessive thirst are the most common symptoms. In children who were previously dry at night, enuresis may be an early symptom of diabetes. The diagnosis is usually straightforward. In chronic renal failure, the total solute load may be normal, but the reduction in the number of nephrons results in a greater than normal load per nephron and consequent polyuria, which is usually of only moderate severity. The polyuria that may follow the relief of chronic urinary tract obstruction is also partly due to this mechanism, but some defect of concentrating power may also be present. Diuresis is also common during recovery from acute tubular necrosis; this is due partly to the elimination of water and electrolytes retained during the phase of oliguria, and partly to incomplete recovery of tubular function.

POLYURIA DUE TO WATER DIURESIS
The simplest cause of this is an increased water intake, which may reach pathological dimensions in the condition known as psychogenic polydipsia or compulsive water drinking. This is a hysterical manifestation and simulates diabetes insipidus. The differentiation is discussed below but, clinically, marked fluctuations in urine output would strongly suggest psychogenic polydipsia. Patients with Sjögren’s syndrome may try to relieve the dryness of their mouths by drinking large volumes of water, with a consequent modest polyuria.

The urine is normally concentrated in the distal tubules and collecting ducts. Antidiuretic hormone (ADH) is secreted by the posterior pituitary in response to a rise in plasma osmolality; its action is to increase the permeability of the tubular epithelium to water. The effect of this is to increase the transport of water from the tubular lumen into the hypertonic renal medulla through which these tubules pass. Thus, a pathological water diuresis may be due either to failure of secretion of ADH or to failure of the renal tubules to respond to its action. Cranial (neurogenic) diabetes insipidus is often caused by an identifiable lesion of the hypothalamus or pituitary (or both), but in about one-third of cases no such cause can be found. Tumours in that region are a common cause and include craniopharyngioma, pinealoma, glioma and metastases from distant primary growths. Diabetes insipidus may also follow trauma to the skull (including operative), infections such as exanthemas in childhood, or infiltration with granulomatous lesions such as sarcoidosis or histiocyotosis X. In such cases, other evidence of hypothalamic pituitary disease may well be present. Occasionally, diabetes insipidus is complicated by a destructive lesion of the ‘thirst’ centre in the hypothalamus so that polydipsia does not accompany the polyuria; water loss is severe, and hypernatraemia with brain damage may result. There is a very rare familial form of cranial diabetes insipidus, inherited as an autosomal dominant trait; even rarer is the DIDMOAD syndrome (diabetes insipidus, diabetes mellitus, optic atrophy and deafness), with autosomal recessive inheritance.

Failure of the renal tubules to respond to the action of ADH is termed nephrogenic diabetes insipidus. A familial form is seen in males only, inherited as a sex-linked recessive (X-linked). It may be a part of other renal tubular defects such as the Fanconi syndrome, with cystinosis and proximal renal tubular acidosis (type 2), and it can also occur in renal amyloidosis, myelomatosis and hyperglobulinaemia.

The differentiation of the two types of diabetes insipidus from each other and from psychogenic polydipsia may be difficult. This is because a prolonged water diuresis from any cause may lead to partial resistance to the action of ADH. Thus, deprivation of water in psychogenic polydipsia may not cause immediate cessation of the polyuria, although there will usually be a considerable reduction. In diabetes insipidus, whether nephrogenic or of cranial origin, the polyuria will continue despite water deprivation, and the patient becomes very thirsty and ill; the test is not without its dangers in this situation. Further information may be obtained from the administration of desmopressin, an ADH analogue. This will clearly have no effect in nephrogenic diabetes insipidus, but it will reduce the urine output in cranial diabetes insipidus and in psychogenic polydipsia. Once again, the result may not be clear-cut, and the effect of desmopressin may not be apparent for several days. There is a danger that, in psychogenic polydipsia, the continued ingestion of large amounts of water after the administration of ADH may cause water intoxication.

Several other conditions cause nephrogenic diabetes insipidus. Polyuria is a common feature of potassium depletion; this condition might be suspected if muscular weakness is a prominent complaint or the
Polypoid swelling: A herniation of the synovial membrane that occurs at an early stage. The polyuria due to the latter must not be confused with the osmotic diuresis of chronic renal failure discussed above.

**TRANSIENT POLYURIA**

Polyuria lasting for only a few hours can occur in various circumstances. It is rarely of any great significance and is often physiological. The diuresis that follows excessive water drinking needs no comment. The same applies to polyuria in the course of diuretic therapy, although the diuretic effect of such substances as tea or coffee may occasionally provoke a complaint from a patient who has not realized the association.

Cold weather may also induce polyuria as a result of reduced fluid loss from the skin; travellers returning from a long stay in the tropics and accustomed to a large fluid intake may occasionally complain of polyuria on return to a colder country. It is this contrast in various circumstances. It is rarely of any great significance and is often physiological. The diuresis that follows excessive water drinking needs no comment. The same applies to polyuria in the course of diuretic therapy, although the diuretic effect of such substances as tea or coffee may occasionally provoke a complaint from a patient who has not realized the association.

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its origin. The ‘cysts’ may be multiple. Such extensions of the knee joint may sometimes rupture and cause an inflammatory reaction in the calf muscles, and this may be mistaken for a deep venous thrombosis. In diagnosis, CT or magnetic resonance imaging may be of value.

**Varicosities veins**

Varicosities of the short (small) saphenous vein are often present in the popliteal space; the diagnosis presents no difficulties, as the veins in the lower part of the leg will also be varicose. However, they become much more obvious when the patient stands. Although the above-described are the most common causes of popliteal swelling, the following conditions are much more rarely encountered.

**Acute abscess**

This is recognized by the signs of acute inflammation; the skin is red and oedematous, the pulse and temperature are raised, and the swelling is very painful. The knee is kept flexed in order to minimize the tension of the part. The abscess may be caused by suppurative lymph nodes or by suppurative periostitis or necrosis of the lower end of the femur. In the former case, the abscess will be superficial; in the latter case, it is deep to the popliteal vessels.

**Aneurysm of the popliteal artery**

This gives rise to an expansile pulsating tumour, the pulsation being synchronous with the heartbeat. Pressure on the femoral artery above will cause a diminution in size of the swelling and cessation of pulsation. The pulse at the ankle on the affected side may be smaller than that on the opposite, and delayed. If a stethoscope is placed over the swelling, a distinct bruit can be heard.

The major complaint of the patient will probably be of pain, which may be referred down the leg if either popliteal nerve is pressed on, or in the site of the swelling if the bone is eroded. Varicosities veins are almost always present on account of pressure on the popliteal vein. Owing to its pulsatile character, an aneurysm is not often mistaken for anything else, but not every swelling that pulsates is an aneurysm. A soft vascular sarcoma growing from the end of the femur may be pulsatile, and a bruit may be heard over it, but the tumour is not as compressible as an aneurysm is, and the effects on the distal pulse are less marked. A radiograph (or, better, a CT scan) will usually settle the question at once. Distinction must also be drawn between a tumour that pulsates and a tumour to which pulsation is communicated. For instance, an abscess or a solid swelling lying over the popliteal artery may appear to pulsate, but the movement is heaving in character and not expansile. In the rare event of an aneurysm having become filled with clot, it might be taken for a solid tumour growing either from the soft parts or from the bone. Finally, the aneurysm may present on the medial side of the lower end of the thigh, anterior to the tendon of the sartorius.

**SOLID SWELLINGS NOT CONNECTED WITH BONE**

**Enlarged nodes**

It is not common to find the popliteal nodes enlarged from any cause. It is possible that they may become infected with pyogenic organisms from a sore on the back of the leg.

**Tumours**

Tumours are rare; they may be either innocent (e.g. *lipoma* and *neurofibroma*) or *sarcomatous*, starting in the connective tissue of the popliteal space, or attached to one of the muscles. The innocent tumours are of long history and are well defined; the malignant lesions are rapidly growing and infiltrating. A lipomatous mass, in the popliteal fossa or on the medial aspect of the knee, is not infrequently present in osteoarthritis of this joint and is part of the general fatty infiltration that gives rise inside the joint to the *lipoma arborescens* of the synovial membrane.

**SOLID SWELLINGS CONNECTED WITH BONE**

In all cases of bony tumour, a radiograph should always be obtained.

**Benign tumours**

*Exostoses* may be found, generally in children and young adults, growing from the region of the epiphyseal cartilage of the femur. Others may occur in other parts of the skeleton, and sometimes several members of the family are similarly affected. The swelling is of slow growth, is well defined and rarely causes any problems. It is most often found at the inner side of the popliteal space. Exostoses may be confused with ossification of the insertion of a tendon or muscle, the adductor longus muscle being the most commonly affected (rider’s bone).

*Osteoclastoma* (see Fig. L.45) is prone to occur in the upper end of the tibia or lower end of the femur, and this may cause an asymmetrical expansion of the cortex presenting in the popliteal fossa. Usually, expansion of the bone can be detected on other aspects and, if the condition is advanced, the shell may be so thin in some places that ‘eggshell cracking’ can be elicited. The radiographic appearances are typical – expansion and thinning of the cortex, an absence of new bone formation, and trabeculation.
Posture, Abnormal

Mark Kinirons

The term ‘posture’ refers either to the position of one particular part of the body, such as an arm or leg, or to the position of the body as a whole. Abnormalities of posture may thus be limited to individual parts of the body or to the appearance of the whole body. In general, posture cannot be considered in isolation as part of a differential diagnosis. Posture is altered in many conditions (muscular, skeletal, neurological, psychological, etc.).

In the conscious patient, disorders of the basal ganglia produce some of the most characteristic abnormalities of posture. Dystonia may affect the body as a whole in the generalized dystonic disorders such as dystonia musculorum deformans. Focal dystonias may produce a local abnormality such as spasmodic torticollis or writer’s cramp. Parkinson’s disease is associated with a very characteristic flexed posture in the late stages. In athetosis and chorea, postural abnormalities may be seen in addition to the involuntary movements. Kyphosis, for example, is increasingly common with advancing age. It can be taken to be due to a degenerative arthritis or osteoporosis. Rarely such a posture is hysterical.

Prenomenstrual Syndrome

Tony Hollingworth

Prenomenstrual syndrome (PMS) can be defined as the cyclical recurrence of psychological, behavioural and physical symptoms during the luteal phase of the menstrual cycle. This is essentially the 2 weeks prior to menstruation, and the symptoms resolve by the end of menstruation. The woman should then be free of symptoms between the end of menstruation and subsequent ovulation. Psychological and somatic disturbances are part of the normal physiology of the menstrual cycle; however, when exaggerated, they may lead to severe psychological disturbance and behavioural abnormalities. The symptoms include bloating, cramping, pain and tenderness in the breasts, a temporary gain in weight, and some swelling of the hands and feet. They can also include emotional tension, bad temper, nervousness, irritability, headache, lack of concentration, depression and insomnia, sufficient to interfere with the normal enjoyment of life. The majority of women (95 per cent) will have some premenstrual symptoms, only a small percentage (5 per cent) being totally symptom-free. In a small group of women (5 per cent), the symptoms of PMS have a major impact on their lives, have led to suicide and acts of aggression, and have even been cited as a defence in murder trials.

The American Psychiatric Association has established guidelines for the diagnosis of PMS, which it has designated as premenstrual dysphoric disorder. These criteria are as follows:

- Symptoms are temporally related to the menstrual cycle during the last 2 weeks of the luteal phase and resolve after the onset of menses.
- The diagnosis requires at least five of the following, and one of the symptoms must be one of the first four:
  - Markedly depressed mood, feelings of hopelessness
  - Marked anxiety or tension
– Marked affective lability, for example a sudden onset of being sad, tearful, irritable or aggressive
– Persistent and marked anger or irritability, or increased interpersonal conflicts
– A decreased interest in usual activities.
– Easy fatigability or a marked lack of energy.
– A subjective sense of difficulty in concentrating
– Changes in appetite, overeating or food craving
– Hypersomnia or insomnia
– A feeling of being overwhelmed or out of control
– Physical symptoms such as breast tenderness, headaches, oedema, joint or muscle pain, or weight gain
• The symptoms interfere with work or usual activities or relationships.
• The symptoms are not an exacerbation of another psychiatric disorder. Thus, PMS is, in large part, a diagnosis of exclusion.

The aetiology remains obscure and consequently presents therapeutic difficulties. The underlying cause may be a combination of imbalances or abnormalities of ovarian steroid production and central nervous transmitters. It has been shown that women with PMS have lowered levels of whole blood serotonin concentrations and platelet serotonin. Several serotonin reuptake inhibitors (SSRIs) have now been shown to improve symptoms of PMS. Elimination of the cyclical ovarian function results in the complete suppression of symptoms. Despite cyclical ovarian steroid function being the trigger for PMS, there is no definitive test to distinguish it from other disorders. The nature of the symptoms is less important than the timing, and keeping a diary of symptoms may be useful to distinguish primary PMS. The latter are a group of patients who have true PMS with underlying psychopathology. If the symptoms do not follow the pattern described, an alternative diagnosis needs to be considered.

Treatment depends on the severity of symptoms and may consist of a mixture of counselling, education, exercise and reassurance in patients with mild problems. Essential fatty acids and pyridoxine have been used in the past. SSRIs may now be used to good effect in moderate cases. In more severe cases, suppression of ovulation can be used, and this may start with continuous combined oral contraception usage, providing there are no contraindications to the use of oral contraceptives. Continuous progestogens, Danazol and gonadotrophin-releasing hormone analogues can be used. If these are unsuccessful, total abdominal hysterectomy and bilateral salpingo-oophorectomy may in severe cases need to be considered, with subsequent hormone replacement therapy in the form of continuous oestrogen.

For further information, go to www.womenshealth.gov/faq/pms

PRIAPISM

Ben Challacombe

(See also PENIS, PAIN IN, p. 506.)

Priapism signifies a persistent, usually painful, erection of the penis that is generally not accompanied by sexual desire. The term priapism is named after the Greek god Priapus, who was cursed with a huge, permanently erect penis. This is a medical emergency, and the longer the time interval between development of the priapism and presentation to a physician, the worse the eventual prognosis is likely to be. Untreated, the penis eventually undergoes fibrosis of the corporeal smooth muscle and cavernosal artery thrombosis, destroying the erectile mechanism. There are two main types of priapism: low flow and high flow. The former is exquisitely painful, with a rigid erection; the latter is not. Differentiation between the two initially involves aspiration of corporal blood and analysis of the level of oxygenation. Low-flow priapisms contain hypoxic blood with a reduced pH, unlike the high-flow variety, which contain arterial levels of oxygenation.

The history should include previous pelvic surgery and pathology, sickle-cell disease, drug use, previous similar episodes and a history of pelvic/perineal trauma. A urologist should be contacted as a matter of urgency.

LOW-FLOW (ISCHAEMIC) PRIAPISM

The important causes are:

• Pharmacological: which can follow illicit or prescription drug use. Increasing numbers of men are using injectable agents (intracavernous injections, e.g. papaverine or alprostadil) to recover erectile function after prostatic and bladder surgery, or as a result of erectile dysfunction that is resistant to oral agents. Oral phosphodiesterase inhibitors, such as sildenafil citrate (Viagra), tadalafil (Cialis) and vardenafil (Levitra), may, if used by men with normal erectile function, also result in priapism. Other groups reported are antihypertensives, antipsychotics (e.g. chlorpromazine and clozapine), antidepressants (most notably trazodone), anticoagulants and recreational drugs (alcohol and cocaine).
PRURITUS ANI

- **Spinal cord injury:** after injury to the upper dorsal region of the spinal cord. The damage may have produced fracture–dislocation of the spine with paraplegia, in which case the diagnosis will be obvious; short of this, however, a minor degree of injury, with contusion and small haemorrhages into the substance of the cord, may be followed by priapism, sometimes persisting for weeks before recovery occurs. Rarely, cerebrospinal syphilis or tumour may also be responsible.

- **Leukaemia:** priapism may be the presenting feature of or be seen during both myeloid and lymphocytic leukaemia. The cause of the priapism in leukaemia is related to sludging of abnormal blood cells within the penile corpora. This diagnosis is suggested by concomitant splenomegaly and/or lymphadenopathy, and it is confirmed by the haematological findings.

- **Sickle-cell disease and trait:** here the priapism is usually related to a concurrent sickle-cell crisis resulting in abnormal erythrocytes blocking the penile vasculature. This type of priapism is often highly resistant to treatment, and it may be preceded by an intermittent or ‘stuttering’ priapism (see below). Thalassaemia may also rarely cause priapism.

- **Rarely, malignancy** of the urethra, either primary or secondary to carcinoma of the prostate, bladder or testis: this is caused by local invasion from the tumour, and these diseases may also cause a high-flow variety.

- **Other very rare causes** include total parenteral nutrition, Fabry’s disease, dialysis, vasculitis, amyloid, malaria and fat embolism.

*Intermittent or stuttering priapism* is the term used to describe frequently repeated prolonged erections that are of long duration but lack the persistence of true priapism. This is particularly common in young men with sickle-cell disease and often precedes a full-blown event. Seldom will priapism be the only symptom in the case; the diagnosis will be made from the history and from the other symptoms and signs.

**HIGH-FLOW (NON-ISCHAEMIC) PRIAPISM**

High-flow priapisms are generally not painful and may not be fully rigid, patients may still be sexually active, a straddle injury is usually the initiating event, and there may be chronic recurrent presentations. This type of priapism is usually not caused by medication, but it often follows pelvic trauma, such as road traffic accidents, and is caused by a fistula between the cavernosal artery and the corpus cavernosum. Occasionally, cocaine use can initiate a high-flow priapism. Diagnosis is usually made at angiography, where selective embolization usually resolves the condition. It can also be caused by more minor perineal trauma following a bicycle accident, and by local invasion of malignant tumours.

**PRURITUS ANI**

Harold Ellis

Pruritus ani – the sensation of itching around the anal verge – is a common symptom. In more than half the patients, no obvious cause can be found (idiopathic pruritus), but in every case the following checklist should be considered:

- The pruritus may be the result of a general disease associated with itching (see PRURITUS, GENERALIZED, p. 527). Examples are lymphoma, advanced renal failure, severe jaundice and diabetes mellitus. The latter is often associated with Candida albicans infection (thrush), and this may occur, of course, in the non-diabetic patient. As Candida is often a secondary invader on any moist and excoriated skin, however, it may well not be the primary cause of the condition.

- The localized itching may be due to a skin disease that happens particularly to affect the perianal region. Examples include: scabies, where characteristic lesions may be seen elsewhere in the body, notably between the fingers and on the anterior aspects of the wrists; pediculosis pubis, where the parasites may be noted in the anal region, as well as in their usual site in the pubic hairs; and fungal infection. The latter is particularly to be thought of where the skin lesion has a well-defined border at its lateral extent. Other lesions may be found between the toes and in the groin, and proof may be obtained by examination of scrapings from the affected skin. Erythrasma, due to infection with Corynebacterium minutissimum, may be diagnosed by demonstrating coral-pink luminescence when viewed with a Wood’s lamp.

- Any condition within the anus or rectum that produces moisture and sogginess of the anal skin is liable to cause pruritus ani. These lesions include prolapsing piles, prolapse of the rectum, anal fissure, anal fistula, anal papillomata or condylomata, carcinoma or benign tumours of the rectum, colitis or colonic Crohn’s disease (Fig. P.20). Anal incontinence due to sphincteric injury may result in constant soiling of the perianal skin. Careful inspection of the anal verge, digital examination of the anal canal, proctoscopy and sigmoidoscopy, where necessary, will rapidly expose the underlying cause of this condition.
Excessive sweating, especially in hot weather and in hairy men, may be associated with pruritus ani, especially in subjects who wear thick and rough undergarments.

Pruritus ani is unusual in children and, when it occurs, a well-recognized cause is infestation with threadworms (Enterobius vermicularis) or Corynebacterium minutissimum. Characteristically, the worms migrate to the anal verge (especially at night), and scratching results in autoinfection. The parasite is white, about 6 mm long and the thickness of cotton thread. The parasites may be noted at the anal verge or seen at proctoscopy. If the diagnosis is suspected but no parasites are immediately seen, a wash-out of the rectum with normal saline should be inspected against a black background, when the white parasites can be detected. It should be noted that threadworm infection may also occur in adults.

Idiopathic pruritus is diagnosed when no obvious cause can be found. A number of theories have been suggested, including: allergy; that the original cause has now disappeared but the pruritus has persisted because of continued scratching of the anal region by the patient; irritation of the perianal skin by faecal contamination even when no gross soiling is evident; or some psychogenic cause.

Figure P.20 Severe pruritus extending forward to the vulva in a young girl with extensive Crohn’s disease of the colon and distal rectum.

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Figure P.20 Severe pruritus extending forward to the vulva in a young girl with extensive Crohn’s disease of the colon and distal rectum.

Figure P.21 Excoriations.
itches, and the intense nocturnal itching of scabies. In general, the most itchy dermatoses — apart from the infestations — are dermatitis of all causes, particularly atopic dermatitis, lichen planus, pruritic heat, urticaria, including cholinergic urticaria and dermographism, dermatitis herpetiformis, pemphigoid gestationis, cutaneous T-cell lymphoma, and onchocerciasis.

Generalized pruritus
- Senile asteatosis
- Liver: obstructive hepatopathy (including primary biliary cirrhosis)
- Kidney: chronic renal failure
- Blood: polycythaemia, anaemia, iron deficiency
- Endocrine: hyperthyroidism, myxoedema
- Malignancy: lymphoma, liver secondaries, T-cell leukaemia
- Pregnancy: last trimester (especially in atopics)
- Hydroxyethyl starch (plasma expanders)
- Psychological: delusional state of parasitosis
- Drugs, gold salts, detergents, degreasing agents

Faced with a scratching patient, it is important not to overlook an infestation (e.g. scabies or pediculosis) (Fig. P.22). One condition often forgotten by Western physicians is onchocerciasis (recently estimated by the World Health Organization to affect 20 million people worldwide), in which pruritus may be intense, and localized to the legs and buttocks or involve the whole body. Onchocerciasis occurs in a broad belt worldwide from 19° North to 15° South, as well as scattered foci in Mexico, Central and South America, the Yemen and parts of Saudi Arabia. Onchocerca nodules containing the adult worm may be found around the pelvic girdle or on the head and neck, and the diagnosis will be confirmed by finding microfilariae from skin snips.

In the elderly, the most common cause of generalized pruritus is a subclinical dryness of skin, senile asteatosis, due to inadequate secretion of sebum and sweat. A subtle scaling may be seen, and perhaps an eczema craquele on the anterior shins or forearms (Figs P.23 and P.24). The rather evanescent but intensely itchy rash of cholinergic urticaria can also be easily missed. Likewise, patients who have received infusions of plasma expanders based on hydroxyethyl starch may experience an intense and persistent itch out of all proportion to the rather minimal rash. If no primary dermatosis can be identified, the following possibilities should be considered:

- **Liver disease:** pruritus is severe in obstructive jaundice, but in hepatitis and primary biliary cirrhosis can begin before icterus is obvious.
- **Renal disease:** the skin in advanced uraemia may be intensely pruritic and, in patients receiving dialysis, sudden shifts in electrolytes may cause pruritus, as may secondary hyperparathyroidism.

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**Box P5 Causes of pruritus**

**Itchy dermatoses**
- Infestations (scabies, lice, insect bites)
- Dermatitis — all causes
- Lichen planus
- Prickly heat
- Urticaria, including cholinergic urticaria and dermographism
- Dermatitis herpetiformis
- Pemphigoid gestationis
- Cutaneous T-cell lymphoma
- Onchocerciasis

**Generalized pruritus**
- Senile asteatosis
- Liver: obstructive hepatopathy (including primary biliary cirrhosis)
- Kidney: chronic renal failure
- Blood: polycythaemia, anaemia, iron deficiency
- Endocrine: hyperthyroidism, myxoedema
- Malignancy: lymphoma, liver secondaries, T-cell leukaemia
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- Psychological: delusional state of parasitosis
- Drugs, gold salts, detergents, degreasing agents

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**Figure P.22** Cutaneous larva migrans.

**Figure P.23** Acute eczema.
• **Blood disease:** pruritus is common in polycythaemia rubra vera and also in iron deficiency, even without anaemia.

• **Endocrine disease:** generalized itching is occasionally a presenting feature of hyperthyroidism, and it can accompany the dry skin of myxoedema. Diabetes mellitus is said to cause generalized pruritus, but this is more often found to relate to localized infection with *Candida* (e.g. candidal vulvo-vaginitis).

• **Malignancy:** generalized pruritus, sometimes with prurigo nodules, can be the presenting feature of underlying cancers, especially lymphomas. T-cell leukaemia – particularly the Sézary syndrome – produces a notoriously itchy erythroderma.

• **Pregnancy:** itching is common during the last trimester of pregnancy, and a proportion of patients, probably atopics, will produce prurigo nodules on the arms and legs. The itch disappears after parturition.

• **Psychological:** scratching may be associated with delusions, particularly delusions that parasites are crawling over the skin – a symptom that may indicate serious underlying mental disturbance. Cocaine is also said to be capable of producing an intense itching sensation that has been likened to ants crawling on the skin (formication). This is a diagnosis that should not be overlooked, and the cause may not be readily admitted by the patient.

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**PRURITUS VULVAE**

**Tony Hollingworth**

Pruritus vulvae is a term used to denote vulval itching, which can be intense, unpleasant, embarrassing and, in some cases, debilitating. It may be due to a generalized skin or systemic condition, or be localized due to a specific vulval dermatological condition. It may also be due to a local vaginal cause (Box P.6).

**GENERAL CAUSES OF PRURITUS VULVAE**

These include:

- Skin diseases, including eczema, psoriasis and contact dermatitis to local deodorant use, washing products or man-made fibres. Intertrigo can occur whenever the skin is moist and susceptible to inflammation and secondary infection. This may be a particular problem in overweight women, especially if they have diabetes, in which case they may have a greater chance of developing a candidal infection.

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**Box P.6 Causes of vulval itching**

<table>
<thead>
<tr>
<th>Generalized disease</th>
<th>Localized (vulval) disease</th>
</tr>
</thead>
<tbody>
<tr>
<td>Infammatory skin disease</td>
<td>Infammatory vulval disease</td>
</tr>
<tr>
<td>Atopic eczema</td>
<td>Lichen sclerosus</td>
</tr>
<tr>
<td>Eczema (other causes, e.g. contact)</td>
<td>Lichen planus</td>
</tr>
<tr>
<td>Psoriasis</td>
<td>Lichen simplex chronicus</td>
</tr>
<tr>
<td>Lichen planus</td>
<td>Seborrhoeic dermatitis</td>
</tr>
<tr>
<td>Pityriasis rosea</td>
<td>Eczema</td>
</tr>
<tr>
<td>Urticaria</td>
<td>Psoriasis</td>
</tr>
<tr>
<td>Xerosis (dry skin)</td>
<td>Vulvodynia</td>
</tr>
<tr>
<td>Hyper/hypothyroid disease</td>
<td>– Plasma-cell vulvitis</td>
</tr>
<tr>
<td>Liver disease</td>
<td>Localized pruritus</td>
</tr>
<tr>
<td>Chronic renal failure</td>
<td>Overwashing/detergents (irritant vulvitis)</td>
</tr>
<tr>
<td>– Haematological malignancy</td>
<td>Atrophic vulvitis</td>
</tr>
<tr>
<td>– Drug rashes</td>
<td>Psychogenic</td>
</tr>
<tr>
<td>– Occult malignancy</td>
<td>Vulval intraepithelial neoplasia</td>
</tr>
<tr>
<td>– Iron deficiency</td>
<td>Sexually transmitted disease</td>
</tr>
<tr>
<td>– Psychogenic causes</td>
<td>– Vulval warts</td>
</tr>
<tr>
<td>Infections and infestations</td>
<td>– Candida</td>
</tr>
<tr>
<td>Scabies</td>
<td>– Pubic lice</td>
</tr>
<tr>
<td>HIV-related skin disease</td>
<td>– HIV, human immunodeficiency virus associated skin disease</td>
</tr>
<tr>
<td>Tinea (fungal skin disease)</td>
<td></td>
</tr>
<tr>
<td>Body and pubic lice</td>
<td></td>
</tr>
</tbody>
</table>

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**Figure P.24** Excoriated eczema.
PRURITUS VULVAE

• Drug reactions including latex condoms
• Psychogenic problems
• Medical conditions, which include:
  – Diabetes
  – Hypothyroidism
  – Liver disease
  – Chronic renal failure
  – Crohn's disease
  – Haematological disorders including polycythaemia, leukaemia and Hodgkin's disease.
  – Iron deficiency may be a cause so, in some women with this symptom, it may be useful to check the serum iron level, which may be low. These women are not necessarily anaemic, but their iron stores are low, and treatment with oral iron to supplement their diet can improve their symptoms.

LOCAL CAUSES OF PRURITUS VULVAE

These include:

• Irritant vulvitis due to overwashing, use of panty liners which can make the skin very dry, and ‘wetwipes’
• Atrophic vulvitis in postmenopausal women
• Vulval dystrophies
  – Lichen sclerosus
  – Lichen planus
  – Lichen simplex chronicus
• Premalignant disease (vulval intraepithelial neoplasia)
• Seborrhoeic dermatitis
• Infections
  – Candida
  – Herpes
  – Warts
  – Threadworm, lice and scabies
  – Other sexually transmitted diseases

Infection of the vagina and vulva with Candida albicans (thrush) is a common problem, which may be associated with combined oral contraceptive use, antibiotics, diabetes and pregnancy. The itching is intense, and is associated with a typical curdy, white discharge that causes local inflammatory change, resulting in a very reddened vagina. It can be treated with local preparations of antifungal agents; fluconazole can be given systemically if local treatment is unsuccessful. Vaginal discharge from any cause may be responsible for pruritus, as may infection from the urinary tract. The mucoid discharge from an ectropion, however, does not usually cause pruritus, and infection with Trichomonas vaginalis is more inclined to give rise to vulval soreness. Pubic lice and scabies can cause considerable distress from itching and can be treated with topical agents.

Vulval dystrophies are a common cause of pruritus vulvae. They include:

• Lichen sclerosus is a well-recognized vulval dermatosis that has typical features of loss of the architecture of the vulva with fusion of the labia majora and minora. The skin appears thickened and porcelain-white, but it can be thin and reddened (erythema) as a result of scratching. The skin can appear like cigarette paper. It is thought to be an autoimmune condition associated with other autoimmune conditions especially in the thyroid gland. Treatment of this condition is with high-dose steroid cream (Dermovate) and there is also a need for an emollient and moisturiser. The Dermovate is weaned down once the symptoms are under control but, if the condition flares up, the same regimen can be repeated. There is a small risk that this may predispose the woman to invasive cancer of the vulva (up to 4 per cent of cases). It should be emphasized that carcinoma of the vulva is an extremely uncommon condition in the UK.
• Lichen planus may affect the vulva in isolation and may be destructive of the vulval architecture as well as being intensely itchy. Differentiating between lichen sclerosus and lichen planus can sometimes be difficult, but checking the rest of the body can be useful, and 30 per cent of women with lichen planus will have buccal mucosal involvement. The epithelial surface may demonstrate the lacy white hyperkeratosis, and the inflammation may have a more purplish appearance (violaceous). It is usually a self-limiting condition and, if there is difficulty in diagnosis, a biopsy is indicated.
• Lichen simplex chronicus is a condition in which the habit of scratching the vulval skin may cause an appearance of persistent vulval eczema. Clinically, the vulval skin is thickened and can be lichenified. Treatment is usually with a steroid cream plus an emollient and moisturizer.
• Seborrhoeic dermatitis causes itching in the vulva and, in many cases, there is little to see on vulval inspection. Many of these women will have been treated with repeated preparations for Candida. However, if the vulva is inspected with good lighting and a colposcope, the skin may appear reddened with fine fissures present. Treatment is with a less potent steroid plus an emollient and moisturizer.

Vulval intraepithelial neoplasia (precancerous changes) and Paget's disease are also found on the vulva. They both cause pruritus and need biopsying for diagnosis. Conservative measures using steroid creams are adopted for dystrophies without cell atypia, but
excision of the affected area has to be undertaken when atypia denotes premalignancy.

**Patient advocate groups**
- Candidiasis: www.candida-society.org.uk
- Eczema: www.eczema.org
- Herpes: www.herpes.org.uk
- HIV: www.tht.org.uk
- Lichen sclerosus: www.lichensclerosus.org
- Psoriasis: www.psoriasis-association.org.uk
- Vulvodynia: www.vulvalpainsociety.org
- VIN: www.macmillan.org.uk

**PTOSIS**

Reginald Daniel

Ptosis is the term applied to drooping of the upper eyelid with inability to elevate it to the full extent (Fig. P.25). The most common form is a congenital defect and, if the pupil is in consequence covered, urgent surgical correction is indicated to prevent amblyopia. The acquired form is usually caused by paralysis of the IIIrd cranial nerve, when it may also be associated with paralysis of other ocular muscles, either external or internal.

In paralysis of the cervical sympathetic nerve, slight ptosis may be associated with diminution in the size of the pupil on the affected side, retraction of the eyeball (enophthalmos), and absence of sweating – Horner’s syndrome (Fig. P.26). Ptosis occurs in myasthenia gravis, and is diagnosed using the Tensilon test, which is performed by the intravenous injection of edrophonium chloride (Tensilon), a drug that temporarily blocks the action of the enzyme acetylcholinesterase. Improvement of the ptosis immediately after the injection provides strong support for the diagnosis of myasthenia gravis.

Ptosis, associated with oedema and infiltration of the lids, is also found in inflammatory disorders of the conjunctiva and upper lids. Gross oedema may occur in angioneurotic oedema. It also follows direct injury of the elevating muscle or its nerve supply following lid laceration or blunt trauma.

Congenital ptosis is often bilateral and associated with smoothness of the upper lids and an absence of all the usual cutaneous folds. The levator palpebrae may be absent or ill-developed, and efforts to open the eye are made by the occipitofrontalis muscle.

In the congenital condition called ‘jaw-winking’, movements of the jaw – especially lateral movements – cause the ptotic lid to rise. The patient is thus seen to ‘wink’ while eating or chewing. The cause appears to be a misdirection of nerve fibres resulting in a faulty innervation of the levator muscle.

The causes of ptosis are listed in Box P.7.

![Figure P.25 Ptosis of left upper lid. [Moorfields Eye Hospital.]](image1)

![Figure P.26 Horner’s syndrome: ptosis and a constricted pupil on the right side. [Moorfields Eye Hospital.]](image2)

**Box P.7 Causes of ptosis**

**Congenital**
- Paresis or maldevelopment of levator muscle, jaw-winking syndrome, blepharophimosis syndrome

**Acquired**
- Traumatic
  - Eyelid laceration, post-surgical (following enucleation or orbital surgery)
- Neurogenic
  - IIIrd nerve palsy (due to trauma, ischaemia, inflammation, neoplasm or aneurysm), Horner’s syndrome, multiple sclerosis, neurosyphilis
- Myogenic
  - Dystrophy myotonica, myasthenia gravis, primary muscular atrophy, chronic progressive external ophthalmoplegia, senility, iatrogenic, chronic ocular inflammation
- Mechanical
  - Lid tumours, cicatricial, inflammation
PTYALISM

Harold Ellis

Ptyalism means excessive secretion of saliva. It is not always easy to determine whether there really is excess, or whether the patient is merely allowing the normal volume of saliva to dribble from the mouth. Thus, the difficulty may be solely that of swallowing the normal secretion, as in bulbar paralysis. There may be both excess of secretion and difficulty in swallowing, as in mercurial stomatitis. In other instances, there is too much secretion but no difficulty in swallowing, as in functional or hysterical ptyalorrhoea. The first step towards ascertaining the cause is to inquire as to any medicine or drug the patient may be taking orally or applying externally.

Mercury was the most important of these when it was used as a drug in the treatment of syphilis; its effects were worst when the mouth was not kept clean. Iodides, bromide and arsenic were also often responsible in the past.

If the salivation is not attributable to any drug, it may be the result of one of the many forms of general stomatitis. The nature of a severe stomatitis will be ascertained by local examination; by bacteriological examination of swabblings from the mouth; by serological tests for syphilis; or by microscopical examination of a fragment of the affected tissues.

Tuberculous stomatitis is one of the rarer but severe forms; it may be primary but is more often associated with pulmonary tuberculosis.

If drugs and general stomatitis can be excluded, a local examination may still serve to detect a cause acting by reflex irritation of the Vth cranial (trigeminal) nerve, especially:

- A jagged carious tooth
- A stump left beneath a dental plate
- A broken or ill-fitting dental plate
- A foreign body impacted in the gum
- An ulcerating tumour of the oral cavity

If appropriate examination serves to exclude these causes, the salivation – apparently rather than actually increased – may be found to result from mechanical difficulties in swallowing (see DYSPHAGIA, p. 142). The excessive salivation seen in many cases of advanced carcinoma of the oesophagus results from the oesophago-salivary reflex; a constant excess flow of saliva is secreted in an attempt to ‘swallow’ the obstructing bolus of tumour in the gullet.

In the absence of an obvious local structural lesion, apparent salivation may be due to inability to swallow, as in cases of:

- Parkinsonism
- Bulbar paralysis
- Pseudobulbar paralysis
- Bilateral facial paralysis
- Myasthenia gravis
- Hypoglossal nerve paralysis

The differential diagnosis of these conditions is discussed elsewhere. It is only in bulbar and pseudobulbar paralysis that the dribbling of much saliva is a prominent symptom. Pseudobulbar paralysis, being of cortical and not of medullary nuclear origin, does not exhibit wasting of the tongue.

Slovenliness and lack of cerebral control are responsible for the slobbering and salivation of some elderly or patients with learning disabilities.

PUBERTY, DELAYED

Paul Carroll

Puberty is described as delayed when a boy or girl has passed the normal age of onset of puberty with no physical or hormonal signs of secondary sexual development. Puberty may be delayed for years and still occur normally, in which case it is termed constitutional delay. Delay of puberty may also occur due to undernutrition, chronic ill-health or defects of the reproductive system (hypogonadism) or the body’s responsiveness to sex hormones. Delayed puberty is more common in boys than in girls, and it may be defined as the total absence of sexual development in a boy over the age of 15 years or in a girl who is more than 14 years. Constitutional delayed puberty accounts for half of the male cases, but it is much less likely to be the cause in a girl, in whom over 80 per cent will have some pathological condition.

For practical purposes, the first sign of puberty in a girl is the appearance of pubic hair or a breast bud; in a boy, enlargement of the testes is the earliest sign, before rugosity of the scrotum or pubic hair growth. Prepubertal testes are less than 2 cm in length. If appropriate examination serves to exclude these causes, the salivation – apparently rather than actually increased – may be found to result from mechanical difficulties in swallowing (see DYSPHAGIA, p. 142). The excessive salivation seen in many cases of advanced carcinoma of the oesophagus results from the oesophago-salivary reflex; a constant excess flow of saliva is secreted in an attempt to ‘swallow’ the obstructing bolus of tumour in the gullet.

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Slovenliness and lack of cerebral control are responsible for the slobbering and salivation of some elderly or patients with learning disabilities.
require treatment. Delayed puberty will fall into
one of three categories: (i) constitutional delayed
puberty, (ii) hypogonadotrophic hypogonadism or
hypothalamic–pituitary abnormalities, and (iii) primary
gonadal failure or hypergonadotrophic hypogonadism.
The causes of delayed puberty are listed in Box P8.

CONSTITUTIONAL DELAYED PUBERTY

Constitutional delayed puberty accounts for half of
the cases in boys and only 16 per cent in girls. There is
often a family history, and there may be an indication
of a recent slowing of growth. This may, in fact, be
the normal prepubertal deceleration of growth. Bone
age correlates better with the time of onset of puberty
than does with chronological age. As the bone age
advances, serum gonadotrophin levels increase.
So long as the penis is not very small and there is
no anosmia, it is reasonable to review these patients
every 6 months, especially if the testosterone response
to human chorionic gonadotrophin is appropriate for
the bone age. If the boy is becoming embarrassed
by his lack of sexual development and if his height is
adequate, one may consider low doses of androgen.
Similarly, oestrogens may be used in girls. Generally
speaking, however, these individuals usually develop
perfectly normally, albeit in their late teens or even in
their early 20s.

HYPOTHALAMIC AND PITUITARY CAUSES

Abnormalities of the hypothalamus and pituitary
account for approximately one-third of cases of
delayed puberty in both girls and boys. The cause
may be a space-occupying lesion, trauma or the
result of infection or granulomatous infiltration
involving the hypothalamus, the pituitary or both. The
hormone defects in these cases tend to be multiple,
so that, in addition to delayed puberty due to lack of
gonadotrophins, there may be short stature due to
growth hormone deficiency, lethargy and weakness
due to lack of adrenocorticotrophic hormone (ACTH),
cold insensitivity due to low thyroid-stimulating
hormone levels, or polydipsia and polyuria due
to deficiency of vasopressin (diabetes insipidus).
However, patients with solitary growth hormone
deficiency (see p. 643) may present not only with
short stature (and often obesity) but also with delayed
puberty, although sexual development usually occurs
normally later. Non-secreting pituitary adenomas and
prolactinomas may cause hypogonadism due to the
secretion of high levels of prolactin. High prolactin
levels may also be produced by lesions that interfere
with the production of prolactin-inhibiting factor
(dopamine) or its delivery to the anterior pituitary.

Hypothalamic lack of the releasing hormone for
gonadotrophins (GnRH) accounts for about 7 per cent
of all cases of delayed puberty. This may be due to a
midline developmental defect, since in some cases
anosmia due to hypoplasia of the olfactory bulbs is
present (Kallmann’s syndrome, with a male:female
ratio of 4:1) and there may be a cleft palate, hare lip
and other congenital abnormalities. These patients tend
to be tall because of their hypogonadism.

Male patients have been reported with isolated
luteinizing hormone deficiency. They are tall with
eunuchoidal skeletal proportions (arm span more than
5 cm greater than height, and lower body segment
[heel–pubis] more than 5 cm greater than upper body

<table>
<thead>
<tr>
<th>Box P8 Causes of delayed puberty</th>
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<tr>
<td><strong>Constitutional Hypothalamic syndromes</strong></td>
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<tr>
<td>• Lack of gonadotrophin-releasing hormone (Kallmann’s syndrome)</td>
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<td>• Laurence–Moon–Biedl syndrome</td>
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<td>• Prader–Willi syndrome</td>
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<td>• Lyndi’s syndrome</td>
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<td><strong>Destructive lesions of hypothalamus and/or pituitary</strong></td>
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<td>• Craniopharyngioma</td>
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<td>• Germinoma</td>
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<td>• Chromophobe adenoma</td>
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<td>• Prolactinoma</td>
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<td>• Optic chiasma glioma</td>
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<td>• Meningioma</td>
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<td>• Langerhans cell histiocytosis</td>
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<td>• Hydrocephalus</td>
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<td>• Trauma</td>
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<td>• Vascular lesions</td>
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<td>• Granulomas</td>
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<tr>
<td>• Infections (tuberculosis)</td>
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<td>• Cranial irradiation</td>
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<td><strong>Isolated pituitary deficiencies</strong></td>
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<td>• Growth hormone</td>
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<td>• Luteinizing hormone (fertile eunuch syndrome)</td>
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<td><strong>Gonadal abnormalities</strong></td>
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<td>• Anorchism</td>
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<td>• Ovarian dysgenesis</td>
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<td>– Turner’s syndrome</td>
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<tr>
<td>– Pure dysgenesis</td>
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<tr>
<td>• Noonan’s syndrome (boys and girls)</td>
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<tr>
<td>• Klinefelter’s syndrome (rarely causes delayed puberty)</td>
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<tr>
<td>• Autoimmune ovarian failure</td>
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<td>• Resistant ovary syndrome</td>
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<tr>
<td><strong>Hormonal</strong></td>
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<tr>
<td>– Masculinizing tumour of the ovary</td>
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<td>– Destructive lesions</td>
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<tr>
<td>– Castration</td>
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<tr>
<td>– Mumps (damage occurs very rarely in children)</td>
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<td>– Tuberculosis</td>
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<tr>
<td>– X-irradiation</td>
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<td>– Cytotoxic therapy</td>
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<tr>
<td><strong>Adrenal disease</strong></td>
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<tr>
<td>• X-linked congenital adrenal hypoplasia</td>
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<td>• Cushing’s syndrome</td>
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<td><strong>Thyroid disease</strong></td>
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<td>• Hypothyroidism</td>
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<td>• Hyperthyroidism</td>
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<tr>
<td><strong>Chronic disease (functional gonadotrophin deficiencies)</strong></td>
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<td>• Anorexia nervosa</td>
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<td>• Malnutrition</td>
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<td>• Tuberculosis</td>
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<td>• Severe uncontrolled diabetes</td>
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<td>• Chronic renal failure</td>
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<td>• Cyanotic congenital heart disease</td>
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<td>• Cystic fibrosis</td>
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<td>• Glucose enteropathy and other malabsorption syndromes</td>
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<td>• Connective tissue diseases</td>
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<td>• Haemoglobinopathies</td>
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<td>• AIDS</td>
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<td><strong>Rigorous physical training Drugs</strong></td>
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<td>– Girls</td>
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<tr>
<td>– Androgens</td>
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<tr>
<td>– Anabolic steroids</td>
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<tr>
<td>– Both sexes</td>
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<tr>
<td>– Excess thyroid hormones</td>
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GONADAL ABNORMALITIES

A primary gonadal abnormality is responsible for over a third of cases of delayed puberty in girls but in little over 5 per cent of boys. The most common cause in girls is Turner’s syndrome (see p. 639), in which ovarian dysgenesis is classically associated with the karyotype 45X0. Unlike most other patients with hypogonadism, the girl is short, and there are usually several physical abnormalities, such as a webbed neck, a low hairline posterolaterally, an increased carrying angle of the elbows, short fourth and/or fifth metatarsals and metacarpals, a shield-shaped chest with widely spaced nipples, pigmented naevi, renal anomalies, coarctation of the aorta, atrial and ventricular defects and aortic stenosis. In about a quarter of patients, not all the cell lines carry the karyotype 45X0. This is mosaicism, and cells may be X0/XX, X0/XXY or X0/XX/XXX. Patients with mosaicism tend to be taller than those with X0 Turner’s syndrome, and the physical features outlined above may be less apparent. The serum gonadotrophins luteinizing hormone (LH) and follicle-stimulating hormone (FSH) will be elevated. Buccal smear examinations will usually show absent Barr bodies, indicating the presence of one X chromosome but, if there is any doubt, full karyotyping should be done as there could be mosaicism. In a small number of cases of ovarian dysgenesis, there are none of the features of Turner’s syndrome, the stature is normal and the appearance female, although hypogonadism is present (‘pure dysgenesis’). Half of the patients have a normal female karyotype, 46XX; in the other half, the karyotype is male, 46XY. Serum gonadotrophin levels are high.

Noonan’s syndrome (see p. 641), which looks superficially like Turner’s syndrome, may occur in both sexes and may present with delayed puberty. In the resistant ovary syndrome, the ovaries fail to respond to gonadotrophins, probably owing to a lack of receptors, and an adolescent girl may present with a lack of pubertal development. In a small proportion of girls suffering from autoimmune disease (especially of the adrenal, thyroid and parathyroid glands), delayed puberty may be due to autoimmune ovarian failure; antibodies against the ovaries can be demonstrated in the serum. In boys, anorchism will cause delayed puberty, but this will usually have been investigated long before the expected age of puberty because of the absence of testes in the scrotum. High levels of LH and FSH in the blood will confirm the diagnosis.

In both sexes, surgical removal of the gonads (castration), for whatever reason, will lead to delayed puberty. Mumps usually only damages the postpubertal gonad, and tuberculosis has become rare in the developed world. Irradiation for the treatment of lymphomas is becoming a more common cause of hypogonadism, particularly in girls. The Leydig cells of the testis seem to be more resistant. Cytotoxic chemotherapy may cause gonadal damage. A masculinizing tumour of the ovary will cause delayed puberty in a girl because of the suppression of gonadotrophins by the excess androgens. Secondary sexual hair and even hirsutism will be present, and there may be clitoromegaly.

ADRENAL DISEASE

In girls, congenital adrenal hyperplasia (see HIRSUTISM, p. 283) or Cushing’s syndrome (see p. 282) may lead to failure of breast development and to primary amenorrhoea because of the production of excess amounts of androgen. Pubic and axillary hair will almost always be present, for the same reason. A boy with either of the above two conditions may be thought to be well developed, but examination of his testes will show them to be small because of suppression of pituitary gonadotrophins by the excess androgens. In Cushing’s syndrome in both sexes, the hyperproduction of cortisol also may have an effect in terms of reducing gonadotrophin secretion.

THYROID DISEASE

Where hypothyroidism is associated with delayed puberty in a girl, it is usually due to autoimmune ovarian failure. Hypothyroidism is, in fact, more often associated with precocious puberty (see p. 535). Hyperthyroidism in a child of either sex may lead to delayed puberty due to suppression of gonadotrophin release.

CHRONIC DISEASE

Chronic debilitating diseases of any type, particularly those causing poor nutrition, may cause delayed puberty by interfering with the hypothalamic control of pituitary gonadotrophins. Anorexia nervosa, occurring in about 1 per cent of all teenage girls in Western society, may be missed as a diagnosis, but a careful
history may reveal abnormal eating habits and fads, and evidence of vomiting or purgation. Fine ‘lanugo’ hair may be seen over the body (see p. 159).

Malnutrition such as marasmus and kwashiorkor will be an obvious cause of delayed puberty in underdeveloped countries. Tuberculosis, chronic renal failure, cyanotic congenital heart disease, connective tissue disease and severe uncontrolled diabetes should present no problem in diagnosis. Patients with cystic fibrosis, in addition to having chronic respiratory infections, may also have evidence of gastrointestinal malabsorption due to pancreatic enzyme deficiencies. Gluten enteropathy, which occurs in approximately 1 in 2000 children, may be more difficult to recognize as a cause of delayed puberty. Bowel symptoms may be very subtle, but the diagnosis should be suspected in a short child who has a protuberant abdomen and scanty subcutaneous fat elsewhere. Low red cell folate and antigliadin and antiendomysial antibodies are useful screening tests, but the definitive investigation is a small bowel biopsy. In gluten enteropathy, the villi are flattened.

Rigorous physical exercise in boys and girls can delay the onset of puberty. This is particularly prominent in young female athletes such as ballet dancers and gymnasts, but it also happens in boys. There is evidence that the LH-releasing hormone pulse generator may be inhibited by endogenous endorphins. With the delay in menarche in females, long-term osteopenia may result from chronic low oestrogen levels.

**DRUGS**

In girls, the administration of either androgens or anabolic steroids will lead to suppression of sexual development due to inhibition of gonadotrophin release. Pubic and axillary hair will, however, usually be present. In both boys and girls, the ingestion of excess thyroid hormones in the treatment of hypothyroidism or thyroid cancer may lead to delayed puberty owing to the inhibition of gonadotrophin release.

**PUBERTY, PRECOCIOUS**

Paul Carroll

Before puberty, blood levels of luteinizing hormone (LH) and follicle-stimulating hormone (FSH) are low and are poorly responsive to administered gonadotrophin releasing hormone. At the time of puberty, gonadotrophin secretion begins to increase and gonadal stimulation occurs, with a consequent rise of testosterone in boys and of oestradiol in girls.

In both sexes, puberty is preceded by an increase in the production of the suprarenal androgens dehydroepiandrosterone and androstenedione. The causes of precocious puberty are listed in Box P.9.

In 95 per cent of girls, the first sign of puberty, which is either breast bud or pubic hair development, appears between the ages of 8.5 and 13 years. Puberty is precocious if either of these two events occurs before the age of 8 in a girl. Menarche occurs in 95 per cent of girls between the ages of 11 and 13 years.

In 95 per cent of boys, the testes begin to enlarge between the ages of 9.5 and 13.5 years (mean 11.6 years), and reach adult size between the ages of 13 and 17 (mean 14.9 years). Pubic hair growth develops after testicular enlargement has begun, so it is rare before the age of 9.5 years. Testicular growth is stimulated by the action of FSH on germinal epithelium, and of LH on Leydig cells. The testosterone produced by the Leydig cells is responsible for the growth of the penis. Puberty is precocious in a boy if any of the above-mentioned changes takes place before his ninth birthday.

Precocious puberty can be either a true puberty or a false pseudo-puberty. In true puberty, the changes proceed in the normal physiological manner, albeit

<table>
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<tr>
<th>Box P.9 Causes of precocious puberty</th>
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<tr>
<td><em><em>True (LHRH</em> dependent)</em>*</td>
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<tr>
<td>Constitutional</td>
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<td>Cerebral</td>
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<td>Trauma</td>
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<td>Cranial irradiation</td>
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<td>Granulomas</td>
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<td>Brain abscess</td>
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<td>Hypothyroidism</td>
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<td>Hydrocephalus</td>
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<td>Encephalitis</td>
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<td>Meningitis</td>
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<td>Tuberous sclerosis</td>
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<td>Neurofibromatosis</td>
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<td>Cerebral tumour</td>
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<td>Pineal tumour</td>
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<tr>
<td>Cranial hypophysembloma</td>
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<td>Hamartoma</td>
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<tr>
<td>McCune-Albright syndrome (polyostotic fibrous dysplasia with cutaneous pigmentation)</td>
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<tr>
<td>Russell–Silver syndrome</td>
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<tr>
<td><strong>False (LHRH-independent)</strong></td>
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<tr>
<td>Gonadotrophin-producing tumours</td>
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<td>Hepatoma</td>
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<td>Hepatoblastoma</td>
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<td><strong>Suprarenal</strong></td>
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<td>Teratoma</td>
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<td>Chorionepithelioma</td>
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<td>Congenital suprarenal hyperplasia</td>
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<td>Tumours</td>
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<td>Cushing’s syndrome</td>
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<td><strong>Testicular</strong></td>
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<td>Leydig cell tumour</td>
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<td>Leydig cell hyperplasia</td>
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<td><strong>Ovarian</strong></td>
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<td>Granulosa cell tumour</td>
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<td>Androblastoma</td>
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<td>Lipoid cell tumour</td>
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<tr>
<td>Chorionepithelioma</td>
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<tr>
<td>Benign ovarian cyst</td>
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<td><strong>Drugs</strong></td>
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<tr>
<td>Androgens</td>
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<td>Anabolic steroids</td>
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<td>Oestrogens</td>
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* LHRH luteinizing hormone-releasing hormone
at an early age. In false puberty, on the other hand, changes such as secondary sexual hair growth and enlargement of the phallus occur because of abnormal androgen production, and gonadal function is in fact inhibited. A rare intermediate variety may be produced by non-pituitary, gonadotrophin-secreting tumours.

**CONSTITUTIONAL PREOCIOUS PUBERTY**

This is the most common cause of precocious puberty, especially in girls, where it accounts for 80 per cent of cases. Early puberty in boys is five times less common than in girls, and in boys a pathological cause (especially neurological) is found in 50 per cent. A family history of precocious puberty may be found, although the majority of cases are sporadic. Ovarian cysts may be detected in girls on ultrasound examination. They should not be confused with granulosa cell tumours.

Other variants of premature puberty may be present. *Premature adrenarche* is characterized by the early growth of the pubic hair and a slightly advanced bone age. There are no other signs of premature sexual development, and full puberty develops at the normal age. *Premature thelarche* refers to the early development of breast tissue, often in the second year of life. It usually regresses spontaneously within 2 years, but it may lead on to a full early puberty.

The important implication for a child who enters early puberty is that, because bone maturation is advanced, final adult stature may be below that predicted for an offspring of that family.

**CEREBRAL**

Hypothyroidism is included under cerebral causes of precocious puberty because it seems likely that the mechanism is via the stimulation by thyrotrophin-releasing hormone of gonadotrophins in addition to thyroid-stimulating hormone and prolactin.

A wide variety of other cerebral conditions can give rise to precocious puberty if they are posterior to the median eminence, including the mammillary bodies, and involve the posterior part of the floor of the third ventricle. *McCune–Albright syndrome* is only tentatively included under cerebral causes of precocious puberty because it has been postulated that there may be inappropriate secretion of hypothalamic releasing hormones in this condition. The syndrome is rare, and it is much more common in girls than in boys. Precocious puberty is usual, but other endocrine abnormalities such as thyrotoxicosis, hyperparathyroidism, acromegaly and Cushing’s syndrome may occur. One of the characteristic features is polyostotic fibrous dysplasia, which may be unilateral. There are pigmented areas on the skin roughly corresponding to the underlying bony lesions.

Patients with Russell–Silver syndrome (see STATURE, SHORT p. 645) may develop early puberty. The cause is unknown.

True precocious puberty can develop in children who are adopted from underdeveloped countries. This is thought to be secondary to correction of malnutrition, generally after 3 years of age. The correction of a virilizing condition can be followed by true precocious puberty due to activation of the hypothalamic–pituitary–gonadotrophin–gonadal axis. This occurs in boys and girls with congenital adrenal hyperplasia who have received treatment with glucocorticoids after the age of 4–8 years.

**Gonadotrophin-producing tumours** are exceedingly rare. They produce testicular enlargement in boys and vaginal bleeding in girls as the first sign and are usually malignant.

**ADRENAL CAUSES OF FALSE PUBERTY**

The secretion of excess androgens by the adrenal gland causes false puberty. The growth of secondary sexual hair is stimulated, and the phallus enlarges. Gonadotrophins, however, are suppressed so the testes remain small in a boy, and periods do not develop in a girl. Suprarenal tumours and Cushing’s syndrome can be distinguished from congenital suprarenal hyperplasia by the presence of raised plasma 17-hydroxyprogesterone and urinary excretion of pregnanetriol in the latter when it is due to a 21-hydroxylase defect, and by normal suppressibility of urinary 17-oxosteroids.

**TESTICULAR CAUSES OF FALSE PUBERTY**

Testicular Leydig cell tumours causing false puberty are exceedingly rare. Occasional cases of Leydig cell hyperplasia have been described. The tumours are usually palpable, but occasionally may be small. However, a useful clue to the presence of a testicular tumour is that the contralateral testis is even smaller because gonadotrophins are suppressed. In most cases, the condition is fully established by the age of 6 years. Precocity is marked: the physique is muscular, body hair growth is considerable, the voice is strikingly gruff and manly, and the penis and prostate are enlarged. In half the cases, there may be grossly overt psychosexual behaviour. Because the tumour is secreting testosterone, high levels of 17-oxosteroids are not always found in the urine, but raised plasma testosterone is diagnostic.
OVARIAN CAUSES OF FALSE PUBERTY

Ovarian tumours account for only a few per cent of cases of precocious puberty in girls. They usually present at about the age of 4 years with irregular vaginal bleeding, breast development and pubic and axillary hair growth. There may be abdominal pain, and the tumour may be palpable. The usual type of tumour producing an endocrine effect is a granulosa cell tumour. This is usually benign, but about one-fifth are malignant.

Androgen-producing tumours of the ovary (androblastomas) cause heterosexual precocious puberty with the development of secondary sexual hair, hirsutism and virilization.

DRUGS

The accidental or deliberate ingestion of oestrogens (commonly the mother’s contraceptive pills) by girls may lead to vaginal withdrawal bleeding and to some breast development. Androgens and anabolic steroids may cause secondary sexual hair growth and phallic enlargement.

PUBIC HAIR, LOSS OF

Paul Carroll

The amount of pubic hair varies from individual to individual, and is less in some races than others, for example in those of oriental origin compared with Caucasians. In the female, secretion of the adrenal androgens androstenedione and dehydroepiandrosterone is responsible for the growth of pubic hair, via their conversion into testosterone. In the male, the secretion of testosterone by the testis and its conversion to dihydrotestosterone brings about pubic hair growth. Bearing these facts in mind, it can be seen that any disease process that brings about pubic hair growth. Bearing these facts in mind, it can be seen that any disease process that might cause secondary sexual hair growth and phallic enlargement.

PULSE, ABNORMAL RATE

Gerry Carr-White

A normal pulse rate varies from about 60 to 100 beats per minute: bradycardia is defined as a rate of less than 60, and tachycardia as a rate of greater than 100 beats per minute. This does not imply that rates outside this range are necessarily abnormal; for example, a rate of 40 or so is quite common in athletes in training, and a rate of over 100 would not necessarily be remarkable in an exceedingly anxious patient.

BRADYCARDIA

Bradydardia may be due either to a slow rate of discharge of the sinoatrial node or to various disorders of impulse formation or conduction. It should not be diagnosed solely from the rate as felt at the radial pulse, as in various conditions such as atrial fibrillation or extrasystoles, only a proportion of the beats may reach the wrist; the true heart rate can then be determined only by auscultation. The causes are summarized in Box P.11.

Sinus bradycardia is not uncommon in otherwise normal individuals, especially during sleep when rates as low as 30–40 beats per minute have been recorded during continuous monitoring. The pulse rate may
also be slow during convalescence after influenza and other fevers. In acute nephritis, bradycardia is probably a reflex result of the acute hypertension. A similar mechanism operates in cases of phaeochromocytoma releasing predominantly noradrenaline (norepinephrine); the paroxysms of hypertension are associated with striking bradycardia, unlike the type of attack due to release of adrenaline. Bradycardia is also a well-recognized, but far from constant, finding in obstructive jaundice and when the intracranial pressure is raised for any reason; in the latter case, the slow rate may be due to direct stimulation of the vagal centre. Myxoedema is another cause of bradycardia that may be profound in myxoedema coma. The pulse is also often very slow in anorexia nervosa. The only valve lesion to be associated with a slow pulse is aortic stenosis; in some severe cases, a rate as low as 50 beats per minute may be found, even in the presence of left ventricular failure. As in myxoedema, the reduced metabolic rate of hypothermia, either accidental or induced, is associated with bradycardia; even in atrial fibrillation, a common complication of hypothermia, the ventricular rate is quite slow. As a transient phenomenon, bradycardia occurs in carotid sinus syncope and in vasovagal attacks in general. Drugs that commonly cause bradycardia include beta-blockers, non-dihydropyridine calcium-channel blockers and antiarrhythmic medications such as amiodarone. Bradycardia may also be found after large doses of cholinergic drugs (e.g. carbachol) and anticholinesterases (e.g. neostigmine) and rarely in acetyl choline esterase inhibitors (used in dementia of Alzheimer’s type). Sinus bradycardia may also be seen in digoxin intoxication.

The sick sinus syndrome is caused by impairment of sinoatrial node function. It is thus characterized by sinus bradycardia (Fig. P.27), sinus arrest, which may cause Stokes–Adams attacks, and sinoatrial block. The latter, like sinus bradycardia, may occur normally during sleep and is a result of failure of transmission of a sinus impulse to the atria; a whole electrocardiographic complex is thus deleted. Higher degrees of heart block are more likely to lead to significant bradycardia and Mobitz type 2 second-degree heart block and third-degree heart block (with complete AV dissociation) often lead to the implantation of a pacemaker. With 2:1 sinoatrial block, the electrocardiogram resembles sinus bradycardia, but exercise or atropine will cause the rate suddenly to double. A well-known association of the sick sinus syndrome is the occurrence of paroxysms of atrial fibrillation, often with a slow ventricular rate, and atrial flutter – the so-called bradycardia–tachycardia syndrome.

Junctional rhythm, originating in the atrioventricular node or the main bundle of His, is a common arrhythmia. It can be due to high levels of digoxin but may occur after myocardial infarction or in healthy individuals; the rate is usually around 60 beats per minute. The atria and ventricles contract simultaneously so that, clinically, the diagnostic feature is a cannon wave in the jugular venous pulse with each beat. In the electrocardiogram, the P wave is often not seen or may be inverted in leads II, III and aVF, and may be just before, incorporated in or just after the QRS complex (Fig. P.28). Junctional rhythm is rarely of any serious significance.

Disorders of the atrioventricular conducting system are an important cause of bradycardia. In first-degree heart block, with prolongation of the P-R interval as the only abnormality, the heart rate is normal. The pulse is slow and regular, however, if second-degree heart block has progressed to 2:1 block. This is the case whether the conduction defect has progressed via Wenckebach...
periods (Mobitz type I) or is the more serious Mobitz type II block. Second degree heart block has to be diagnosed from the electrocardiogram (Fig. P.29).

In complete heart block, atrioventricular dissociation is present, and the ventricular rate is that of a pacemaker somewhere in the conducting system distal to the block. The site of this pacemaker can be determined approximately from the surface electrocardiogram. If the QRS complexes are of normal configuration, the pacemaker must be above the bifurcation of the main bundle; a pacemaker in one or other bundle branch produces complexes with the pattern of bundle-branch block. The most common cause of heart block is fibrosis of the atrioventricular bundle and its branches; the cause of this fibrosis is often not known, and the remaining myocardium is usually healthy. Increasingly it is being recognized that genetic causes are commoner than previously realized. In younger patients with complete heart block, genetic abnormalities are the underlying cause in up to 30 per cent of cases (e.g. LAMIN AC and SCN5A mutations which are also associated with cardiomyopathies). Ischaemia – particularly myocardial infarction – is also a common cause, as are digoxin intoxication and cardiomyopathy of almost any type; cardiac amyloidosis is particularly likely to be associated with conduction defects. Even less common causes include myocarditis, particularly diphtheritic and, in South America, trypanosomiasis (Chagas disease); calcification of the atrioventricular rings in and around the aortic valve may encroach on the bundle and cause heart block. Congenital heart block is a rare condition; the ventricular rate is usually rather faster than in the acquired variety.

Figure P.28 Electrocardiogram of a woman, aged 66 years, at 24 hours after inferior myocardial infarction. Apart from the changes of recent infarction, the P wave is inverted in leads II and III and closely precedes the QRS complex. Junctional rhythm at a rate of 60 per minute.
The most common symptom of heart block is the Stokes–Adams attack, particularly if the degree of block is changing. Established complete heart block may also cause some reduction in exercise tolerance with fatigue, dyspnoea and even heart failure. The diagnosis of Stokes–Adams attacks is discussed under FAINTS (see p. 195). Clinical diagnosis of complete heart block is almost always possible. Apart from marked bradycardia with a ventricular rate around 30 or 40 beats per minute or less, the diagnostic signs include 'a' waves in the jugular venous pulse at a faster rate than the arterial pulse with, in addition, cannon waves occurring whenever atrial and ventricular systole happen to coincide. On auscultation, the first heart sound varies markedly in intensity, the louder sounds occurring when ventricular systole follows closely upon atrial systole, so that the atrioventricular valve cusps are wide apart when the ventricles contract (Fig. P.30) (see also HEART SOUNDS, p. 273). Occasionally, it may be possible to hear separate atrial sounds. Importantly, when complete heart block is complicated by atrial fibrillation, none of these signs – which result from a coordinated atrial contraction – is present. An ejection systolic murmur is commonly present due to the large stroke volume, which is also the reason for the wide pulse pressure; these two signs are present in bradycardia from any cause. In the electrocardiogram, P waves and QRS complexes can be identified with no mathematical relationship between the atrial and ventricular rates.

TACHYCARDIA

Tachycardia may be due to an increased frequency of discharge of the sinoatrial node (sinus tachycardia) or to an arrhythmia. (For a discussion of arrhythmias causing an irregular pulse, see PULSE, RHYTHM OF, p. 544.) In this section, only those arrhythmias producing a rapid regular pulse will be discussed in any detail. The common causes of tachycardia are summarized in Box P.12.

<table>
<thead>
<tr>
<th>Sinus tachycardia</th>
<th>Supraventricular tachycardia</th>
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<tbody>
<tr>
<td>Anxiety</td>
<td>Atrioventricular re-entrant tachycardia</td>
</tr>
<tr>
<td>Febrile conditions</td>
<td>Atrial and junctional tachycardia</td>
</tr>
<tr>
<td>Hyperthyroidism</td>
<td>Atrial flutter</td>
</tr>
<tr>
<td>Drugs (e.g. sympathomimetic agents)</td>
<td>Atrial fibrillation</td>
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<tr>
<td>Reflex (e.g. heart failure, hypertension)</td>
<td>Ventricular tachycardia</td>
</tr>
</tbody>
</table>

Sinus tachycardia is present in most febrile conditions due to a direct effect of the pyrexia on the sinoatrial node. In tetanus, the rapid pulse is probably due to involvement of the autonomic nervous system by the toxin. In a few infections, such
as typhoid, the rise in pulse rate may be rather less than expected from the degree of fever; this relative bradycardia is of modest diagnostic significance. The sinoatrial node is affected directly in a number of other situations in which tachycardia is prominent. These include: hyperthyroidism in which there is also the reflex effect of the high cardiac output discussed below; phaeochromocytoma secreting predominantly adrenaline; and anxiety states and other conditions (e.g. severe pain) in which there is a raised level of circulating catecholamines.

Da Costa’s syndrome, known also as cardiac neurosis and effort syndrome, is an important cause of moderate tachycardia that persists sometimes for many years. Various drugs, including atropine and sympathomimetic agents such as ephedrine and isoprenaline, also produce tachycardia, which is in addition a characteristic feature of poisoning by tricyclic antidepressant drugs.

Sinus tachycardia can also be caused by reflex mechanisms. In conditions causing a rise in right atrial pressure, the sinus rate is increased via the Bainbridge reflex. Hypotension from any cause, or more precisely a fall in pulse pressure, also causes tachycardia via baroceptors in the aortic arch and carotid sinus. This is the mechanism, for example, of the tachycardia during the straining period of a Valsalva manoeuvre. In high-output states, the tachycardia is probably secondary to the tendency of the right atrial pressure to rise as a result of the increased venous return; therefore, the pulse rate is rapid in hyperthyroidism, severe anaemia, pregnancy, beri-beri, widespread Paget’s disease and arteriovenous fistula. If such a fistula is accessible, the effect of the high cardiac output can be convincingly demonstrated by the abrupt fall in pulse rate that results from digital occlusion of the fistula. In cardiac failure, the tendency to a reduction in pulse pressure, acting on the aortic and carotid baroceptors, combines with the rise in right atrial pressure to produce considerable tachycardia in most, although not all, cases. Tachycardia is also a feature of severe myocardial disease even in the absence of frank failure; thus, it is found in most cases of myocarditis and in some cases of ischaemic heart disease.

Hypotension due to extracardiac factors is also a cause of tachycardia, which is seen in shock and as a result of the administration of vasodilating agents.

Disproportionate tachycardia on exertion, with a normal resting pulse rate, is seen in patients less severely affected by the conditions discussed above, and is also a measure of an individual’s lack of training. Physical fitness can be roughly quantified by the amount of work that can be done at any given heart rate.

Lesions of the vagus nerve may occasionally cause tachycardia, which has been described with subventricular tumours and in various types of peripheral neuropathy including alcoholic and diphtheritic. There is also a rare primary disorder of the sinoatrial node (sino node re-entry) in which tachycardia is a feature.

There are several varieties of supraventricular tachycardia and these are split into regular and irregular supraventricular arrhythmias. With these rhythm disturbances, the QRS complex will be narrow unless there is associated bundle-branch block. Regular narrow complex supraventricular arrhythmias are due to the following conditions. Atrial-ventricular node re-entrant tachycardias (AVNRTs) are due to re-entry circuits within the AV node (so-called dual AV node pathology) and are distinct from AV re-entrant tachycardia (AVRT), which occurs when there is a conducting pathway between the atria and ventricles in addition to the atrioventricular node, as in the pre-excitation syndromes. In both these conditions, differences in the conductivity and refractoriness of these two pathways can result in a circus movement, with an impulse re-entering the atria from the ventricles via one pathway and then being transmitted again to the ventricles via the other.

There is often no other evidence of heart disease, and paroxysms of tachycardia occur throughout life. The attacks begin abruptly, and patients may describe symptoms of palpitation and light-headedness. The associated anxiety may cause hyperventilation and paraesthesiae in the fingers, or even frank tetany may occur. In such cases, it is clearly important to distinguish cause from effect in the manifestations. The attack may last for minutes, hours or (rarely) days, and it ends as abruptly as it began. Prolonged attacks, lasting for a week or more at a very fast rate, can cause cardiac failure even in otherwise normal individuals, but this resolves rapidly and completely once the attack is over. In older patients, more serious symptoms may occur; in particular, ischaemic pain is common even though the associated coronary artery disease may be quite mild. Paroxysms of AVRT are common in the pre-excitation syndromes, of which Wolf–Parkinson–White syndrome is the most common. In this syndrome, the P-R interval is pathologically short, and the initial part of the QRS complex rises or falls slowly to form the so-called ‘delta’ wave (Fig. P31). In paroxysms of tachycardia in such cases, the QRS complex assumes a normal configuration. This condition is often benign, but paroxysms of atrial fibrillation with a very rapid
Ventricular rate can occur, and sudden death, due probably to ventricular fibrillation, has been reported. Atrial and junctional tachycardia due to enhanced automaticity of a focus in the atria or atrioventricular junction are less common. They may occur in paroxysms but are more often sustained for long periods. There is often some degree of atrioventricular block, which never occurs in AVRT, and there is more frequently associated heart disease such as ischaemia or cardiomyopathy. Digoxin toxicity may also cause such arrhythmias.

Faster atrial rates than those found in the tachycardias already considered produce atrial flutter. Here, the atrial rate is often around 300 per minute; this is faster than the AV node can conduct, and some degree of atrioventricular rhythm at about 150 per minute. This particular rate is rather characteristic of atrial flutter, as it is faster than most sinus tachycardias and slower than many other supraventricular tachycardias. The diagnosis can be made with near-certainty by studying the effect of pressure on the carotid sinus. In atrial flutter, the degree of atrioventricular block is increased, with, characteristically, an abrupt halving of the pulse rate. In sinus tachycardia, carotid sinus pressure produces a more gradual slowing; in other supraventricular tachycardias, the attack is either terminated or continues unabated. In atrial flutter, the electrocardiogram shows rapid regular flutter waves with QRS complexes at one-half or one-quarter of the atrial rate. The atrial rate is so rapid that, in most leads, the flutter waves produce a continuous ‘saw-tooth’ appearance of the baseline (Fig. P32). The absence of any isoelectric segments in the baseline has been suggested as a diagnostic criterion of flutter. There is, however, no real justification for this view, and many authorities prefer to use the term ‘flutter’ for all supraventricular arrhythmias with an atrial rate of 250–350 per minute. The causes of atrial flutter are similar to those of atrial fibrillation (see p. 544), except that it is not so common in mitral valve disease or hyperthyroidism.

Atrial fibrillation presents with an irregular tachycardia, and is discussed at length in the section on PULSE, RHYTHM OF (see p. 544).

The electrocardiographic diagnosis of a supraventricular tachycardia depends on finding a regular tachycardia with P waves that are abnormal both in their timing in relationship to the QRS complex, and in their shape (Fig. P33). The QRS complexes are usually normal, although they may be widened and deformed as a result of aberrant ventricular conduction and resemble bundle-branch block.

Ventricular tachycardia is a much more serious arrhythmia than its supraventricular counterparts. Although it can occur in patients with otherwise normal hearts (often from a focus in the right ventricular outflow tract), there is usually serious underlying heart disease. Much the most common cause is ischaemic heart disease, and this is particularly common after myocardial infarction, when it can cause serious deterioration in the patient’s condition and may be a forerunner of ventricular fibrillation. Ventricular tachycardia is also commonly seen in inherited heart muscle diseases such as the cardiomyopathies (hypertrophic, dilated and right ventricular) and inherited ion channel diseases (such as long QT syndrome, catecholaminergic polymorphic ventricular tachycardia and Brugada syndrome) and in these cases accurate diagnosis and family screening is vital. Q-T prolongation is also commonly caused by drugs including, many antiarrhythmic...
and antipsychotic medications, as well as a number of antihistamines and antimicrobials. In ventricular tachycardia, the rate is very variable (most commonly around 180 per minute, although it may be as low as 100 or less), in which case some prefer to use the term ‘idioventricular rhythm’ rather than ‘tachycardia’. The electrocardiogram shows broad, often notched, QRS complexes with no preceding P waves; a variation in which the QRS axis changes gradually and repeatedly so that the complexes appear to twist around the baseline is known as torsades de pointes. In the usual type, it may sometimes be possible to identify P waves separately at a slower rate; in this case, the diagnosis of ventricular tachycardia is certain (Fig. P34). Otherwise, there can be confusion with supraventricular tachycardia with aberrant ventricular conduction. This is a difficult diagnostic problem as the certain identification of P waves in such a record may be impossible, but it is always safest to treat any regular broad complex tachycardia as ventricular tachycardia until proven otherwise. See also PULSE, RHYTHM OF (p. 544).

Figure P32 Atrial flutter with variable atrioventricular block. The baseline demonstrates rapid, regular flutter waves with characteristic ‘saw-tooth’ appearance.

Figure P33 Supraventricular tachycardia at 160 per minute, probably junctional as P waves cannot be certainly identified, in a patient 1 week after an inferior myocardial infarction.
An essential part of the cardiovascular system examination is assessment of the pulse character, along with the volume, rate and rhythm. The rate and rhythm can be assessed using the radial artery but the character and volume need to be assessed from either the carotid or brachial artery. The character of the pulse is understood to be the shape that would be inscribed by an instrument recording the movement of the artery. The most important deviation from the normal character is in the rate of rise of the pulse. This, together with variations in the amplitude, often referred to as the volume of the pulse, produces patterns that are characteristic of various cardiac lesions – usually of the valves. These observations are best described in simple, unambiguous terms; for example, the pulse volume should be referred to as ‘small/reduced’, ‘normal’ or ‘large/increased’, and not as ‘good’ or ‘poor’.

A common abnormality in the character of the pulse is a rapid rate of rise. The most typical form of this is the pulse of aortic regurgitation. This pulse is of large volume and is described as collapsing, although it also used to be known as a water-hammer pulse. As well as feeling the rapid rate of rise and large volume at the carotid or brachial pulse, you can also examine by feeling with the palm (rather than the tips of the fingers) placed over the wrist, and then elevating the patient’s arm rather sharply above shoulder level.

A collapsing pulse is also found in patients with a large persistent ductus arteriosus or arteriovenous fistula. Other rarer congenital causes include pulmonary atresia with a large ventricular septal defect, and persistent truncus arteriosus. A large volume pulse that is not necessarily collapsing in nature can arise due to high-output cardiac states (thyrotoxicosis, Paget’s disease and severe anaemia), bradycardia and arteriosclerotic arteries in older patients.

Normal or small-volume pulses with a rapid rate of rise are felt in such conditions as gross mitral regurgitation, hypertrophic obstructive cardiomyopathy and fixed subvalvar aortic stenosis. This type of pulse is sometimes described as jerky.

A slow-rising pulse is seen with significant aortic valve stenosis. The volume of the pulse is usually small, and this type of pulse is more easily missed than is the collapsing pulse. Sometimes a notch is felt low down on the upstroke of the pulse, so that there appears to be a very small impulse followed by a much larger one. If this is the case, the pulse is described as anacrotic (an abbreviation of anadicrotic meaning ‘twice-beating on the upstroke’). This pulse can often be felt in the radial and brachial arteries, but it is much more easily detected in the carotids, where it is often accompanied by a systolic thrill – the palpable counterpart of the murmur transmitted to the neck. A variation of this pulse is when the notch is very much higher on the upstroke, producing the impression of two more nearly equal impulses. This is the bisferiens pulse, and it is felt when aortic stenosis is accompanied by aortic regurgitation of at least moderate severity. This type of pulse is rather uncommon and must not be confused with one in which a small notch is felt at the very apex of the pulse; this is sometimes felt in aortic regurgitation, especially if the palpating finger is applied more firmly than usual. A very pronounced bisferiens pulse is very occasionally visible in the carotid arteries.

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An absolutely regular pulse is rare. Even in normal sinus rhythm, slight fluctuations in rate can be detected by measuring successive R-R intervals in the electrocardiogram. This can hardly be detected by palpation of the pulse, however, and this section will deal with irregularities that are apparent clinically. When examining the pulse, it is necessary first to determine whether it is regular or irregular and, if the latter, whether any pattern can be detected within the irregularity. Complete clinical analysis of an irregular
pulsation includes inspection of the jugular venous pulse and auscultation at the apex beat, in addition to feeling the arterial pulse. The venous pulse provides evidence of the presence and frequency of atrial contractions; auscultation allows the detection of ventricular contractions too feeble to produce a pulse at the wrist. Many arrhythmias can be diagnosed clinically, but electrocardiographic confirmation is essential.

There are two main mechanisms that can produce an irregular pulse. The first is a disorder of impulse formation in which, most commonly, an abnormal (ectopic) pacemaker drives the ventricles either directly or via the atrioventricular bundle. The other mechanism is a disorder of conduction in which a block develops somewhere along the long pathway from the sinoatrial node to the ventricular myocardium; if this block is intermittent, an irregular pulse is likely to result. The common causes of an irregular pulse are summarized in Box P.13. For a more detailed discussion of the genesis of arrhythmias, see PULSE, ABNORMAL RATE (p. 537).

**DISORDERS OF IMPULSE FORMATION**

The rate of discharge of the sinoatrial node can vary, more or less rhythmically, in many normal subjects; this condition, termed sinus arrhythmia, is quite benign and hardly justifies classification as a ‘disorder’. The fluctuations in rate are most commonly in phase with respiration, the rate increasing during inspiration and slowing during expiration. This is very common in children and is due to reflex variations in vagal tone (Fig. P.35). The absence of respiratory sinus arrhythmia is sometimes of some slight diagnostic significance in a child, as this is a feature of a large atrial septal defect. There are also two, much rarer, types of non-respiratory sinus arrhythmia. In one, the cycles of tachycardia and bradycardia are much longer and are quite unrelated to respiration; the P waves of the electrocardiogram are normal and constant in configuration. In the other type, the P waves vary slightly in shape, and the irregularity is believed to be due to changes in the site of the pacemaker within the sinoatrial node itself. The irregularity in respiratory sinus arrhythmia is exaggerated by deep breathing and, in all types, is abolished or markedly reduced by exercise or other causes of tachycardia.

**Ectopic beats** are a very common cause of an irregular pulse. They are due to the discharge of an abnormal pacemaker in the atria, atrioventricular junction (atrioventricular node and main bundle of His) or ventricles. Apart from their site of origin, they can be classified as escape beats, extrasystoles and parasystoles. **Escape beats** should be regarded as a protective mechanism against asystole. They arise either from the atrioventricular junction or from the ventricles, and occur whenever, for any reason, there is a prolonged pause in the activity of the sinoatrial node. Thus, they may be seen in sinus bradycardia and during the slow phase of sinus arrhythmia. They are difficult to recognize clinically, but they can be identified in the electrocardiogram by the abnormally long pause preceding an ectopic beat identified, as described below, as junctional or ventricular in origin.

**Extrasystoles** most frequently arise from the ventricles, but atrial and junctional extrasystoles are also common. If a ventricular extrasystole occurs early in diastole, ventricular filling will be incomplete, and the contraction may fail to open the aortic valve; even if it does so, the pulse may be so feeble that it does not reach the wrist. Thus, in the radial pulse, a ‘dropped beat’ will be noticed; on auscultation, the extrasystole will be heard either as the first sound only or as both sounds. Ventricular extrasystoles occurring later in diastole are more likely to produce a palpable impulse at the radial pulse. Often an extrasystole follows each sinus beat to produce pulsus bigeminus.

**Box P.13** Causes of an irregular pulse

<table>
<thead>
<tr>
<th>Disorders of impulse formation</th>
<th>Disorders of conduction</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sinus arrhythmia</td>
<td>Sinoatrial block</td>
</tr>
<tr>
<td>Ectopic beats (supraventricular</td>
<td>Partial atrioventricular block</td>
</tr>
<tr>
<td>or ventricular)</td>
<td>– Mobitz Type I (Wenckebach)</td>
</tr>
<tr>
<td>– Escape beats</td>
<td>– Mobitz Type II</td>
</tr>
<tr>
<td>– Extrasystoles</td>
<td>Apparent irregularity in normal rhythm</td>
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<tr>
<td>– Parasystole</td>
<td>– Pulsus alternans</td>
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<tr>
<td>– Atrial fibrillation</td>
<td>– Pulsus paradoxus</td>
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<tr>
<td>– Atrial flutter with varying</td>
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<tr>
<td>atrioventricular block</td>
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</tbody>
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**Figure P.35** Electrocardiogram showing gross respiratory sinus arrhythmia. The cycle lengths vary from 0.7 to 1.24 seconds.
or coupled beats, or every second sinus beat to produce trigeminus. The patient with extrasystoles may have noticed no abnormality or may describe symptoms of palpitations.

In the electrocardiogram, ventricular extrasystoles are identified by the bizarre configuration of the QRS complex and by the absence of a P wave (Fig. P.36). The pause following the extrasystole is usually fully compensatory; that is, the sum of the R-R intervals preceding and following the ectopic beat equals two complete cycles. Rarely, if the sinus rate is slow, interpolated ventricular extrasystoles occur – the only true ‘extra’ systole. Occasional ventricular extrasystoles are usually benign and do not necessarily indicate organic heart disease; they are more common following overindulgence in tea, coffee, alcohol or tobacco. Frequent ventricular extrasystoles occurring at rest are of more importance and probably indicate heart disease of some kind; however, their exact significance is uncertain and, provided that the standard investigations (including an exercise electrocardiogram) are normal, a good prognosis can be given. They are certainly of more serious significance following myocardial infarction, when, if they are multifocal, or appear in salvos, or are so premature as to deform the T wave of the preceding complex (R-on-T), they may presage ventricular tachycardia or fibrillation.

Atrial and junctional (supraventricular) extrasystoles produce very much the same symptoms and signs as ventricular ones. The electrocardiogram shows abnormal P waves indicating the site of the ectopic focus; with junctional extrasystoles, the P wave is typically inverted in leads II, III and aVF. The form of the QRS complex is usually normal, as the impulse reaches the ventricles via the normal conducting pathways. Occasionally, however, if the supraventricular extrasystole is very premature, one or other branch of the bundle may still be partially refractory so that intraventricular conduction proceeds abnormally and the configuration of the QRS is bizarre, simulating bundle-branch block. This phenomenon is known as ‘aberrant ventricular conduction’ and can only be decisively distinguished from a ventricular extrasystole by the finding of an ectopic P wave preceding the QRS complex (Fig. P.37). The pause following a supraventricular extrasystole is usually less than fully compensatory. Supraventricular extrasystoles are nearly always benign, but if they are frequent – for example in a patient with mitral stenosis – they may indicate that atrial fibrillation is impending.

With extrasystoles in general, the interval between a sinus beat and an extrasystole – the coupling interval – is remarkably constant, implying that the discharge of the ectopic focus is in some way dependent on the preceding sinus beat. This is not the case in the phenomenon known as parasystole. In this situation, an ectopic focus, most often in the ventricles but occasionally in the atria or atrioventricular junction, discharges at its own intrinsic rate regardless of the rate of the sinoatrial node or other dominant pacemaker. The ectopic focus is ‘protected’ from discharge by the normal sinus beats so that, whenever the ectopic discharge finds the ventricles in a non-refractory state, an ectopic beat appears. The diagnosis is made from the electrocardiogram by finding ectopic beats with varying coupling intervals and a succession of interectopic intervals all of which are multiples of a single shorter interval – the intrinsic cycle length of the ectopic focus (Fig. P.38).

The clinical diagnosis of extrasystoles is usually easy but, if they occur frequently, they may be difficult to distinguish from atrial fibrillation. Exercise will usually abolish extrasystoles and, if anything, cause greater irregularity in atrial fibrillation. The jugular venous pulse may also be helpful as cannon waves are
a constant finding in junctional extrasystoles and may occur in other varieties also, if atrial and ventricular systole happen to coincide. A cannon wave implies an effective atrial contraction and therefore rules out atrial fibrillation.

Progressively more rapid rates of discharge of an ectopic atrial focus lead to atrial tachycardia and atrial flutter, which can be regarded as a series of atrial ectopic beats. In these conditions, the ventricular rhythm is usually regular, but when the rate of discharge of the ectopic focus exceeds about 400 per minute, coordinated atrial depolarization and contraction are impossible, and atrial fibrillation results. The supraventricular impulses impinge more or less at random on the atioventricular node, finding it and the remainder of the conducting tissue more or less refractory at any given time so that the ventricular response is totally irregular.

Atrial fibrillation is an extremely common finding in almost any type of heart disease. Probably the most common cause is rheumatic heart disease, especially mitral valve disease; it is much less common, except as a terminal event, in isolated aortic valve disease. It is also a well-known complication in hyperthyroidism, especially in the toxic nodular goitre of older subjects. It is not very common in uncomplicated angina, but it occurs quite frequently after myocardial infarction and when cardiac failure develops. In hypertensive heart
disease, as well, it is rather unusual except late in the course of the disease. Other conditions characterized by chronic atrial fibrillation (as distinct from the transient type discussed below) are constrictive pericarditis, invasion of the pericardium by bronchial carcinoma or other mediastinal tumours, tumours of the heart itself, many varieties of cardiomyopathy and atrial septal defect, but not other types of congenital heart disease. Idiopathic or ‘lone’ atrial fibrillation, with no other evidence of heart disease, is well recognized and not uncommon. Infections – particularly respiratory – can precipitate atrial fibrillation, especially in patients predisposed by rheumatic heart disease. Once the infection is over, sinus rhythm may be restored, either spontaneously or by DC cardio-version. A return to sinus rhythm is also possible if atrial fibrillation has been caused by myocardial infarction, and is almost the rule in thyrotoxic atrial fibrillation once the patient is euthyroid. Other causes of transient atrial fibrillation include pulmonary embolism (although it is rather unusual in chronic cor pulmonale), sick sinus syndrome, alcohol abuse, thoracotomy for any purpose, electric shock and hypothermia, either accidental or induced. It can rarely be caused by drugs such as digoxin and anaesthetic agents.

The development of atrial fibrillation will usually cause the patient to complain of palpitation, and cardiac failure may be precipitated in those with severe heart disease. This is particularly the case in mitral stenosis in which the rapid ventricular rate, by restricting the time available for ventricular filling, causes a steep rise in left atrial pressure and may precipitate pulmonary oedema (Fig. P.39). Once the ventricular rate has been brought under control with antiarrhythmic therapy, most patients with atrial fibrillation have few, if any, symptoms attributable to the arrhythmia. The pulse is totally irregular and rapid in uncontrolled atrial fibrillation. The true ventricular rate can be determined only by auscultation as many of the impulses fail to reach the radial pulse. The difference between the rates as determined at the radial pulse and by auscultation – the ‘pulse deficit’ – is some measure of the lack of control. Once the ventricular rate is under control, there is no pulse deficit, and the irregularity of the pulse, although still present, may be less easy to detect. The other diagnostic feature is the absence of evidence of atrial systole; no ‘a’ waves are seen in the jugular venous pulse and the presystolic murmur of mitral stenosis disappears, as does an atrial gallop rhythm if one has previously been heard. The electrocardiogram shows three diagnostic features: a totally irregular ventricular rhythm; absence of waves; and their replacement by ‘f’ waves, fairly large in amplitude if the atrial fibrillation is of recent onset and becoming smaller as the months and years go by (Fig. P.40).

Atrial flutter, although usually associated with a regular ventricular rhythm, may produce an irregular pulse if the degree of atrioventricular block is variable.

**Figure P.39** Atrial fibrillation with a ventricular rate of 140 per minute, recorded from a man, aged 51 years, with mitral stenosis who was in pulmonary oedema at the time. He became virtually symptom-free when the ventricular rate was controlled with digoxin.

**Figure P.40** Electrocardiogram showing atrial fibrillation with a ventricular rate of 60 per minute. The ‘f’ waves are well shown, as is usually the case in lead VI.
Clinically, this is very difficult to distinguish from atrial fibrillation and, indeed, at fast atrial rates the two conditions merge in so-called ‘flutter-fibrillation’. This unsatisfactory term should be avoided if possible; if each of the varying R-R intervals is a multiple of the interval between two ‘f’ waves, flutter should be diagnosed. For further discussion on atrial flutter, see PULSE, ABNORMAL RATE (p. 537).

DISORDERS OF CONDUCTION

Failure of transmission of the impulse from the sinoatrial node to the ventricles causes ‘dropped beats’. The ventricular rhythm will be regular only if every other beat is dropped (as in 2:1 atrioventricular block) or if the beat is complete and a lower pacemaker is driving the ventricles. Any other pattern of failure of conduction will cause an irregular pulse.

Sinoatrial block is a rather rare conduction defect in which the impulse fails to pass from the sinoatrial node to the atrial myocardium. Although sinoatrial block can be classified into first, second or third degree, only second degree can be diagnosed reliably from the electrocardiogram (Fig. P41). The patients may be symptom-free, but Stokes–Adams attacks can occur. The condition can occur in the absence of other evidence of heart disease, especially during sleep, but it is quite often associated with ischaemic heart disease or, as a transient phenomenon, with acute rheumatic carditis. It is one of the typical features of the sick sinus syndrome, and it can also occasionally be produced by digoxin.

Partial atrioventricular block can also produce an irregular pulse. In Mobitz type 1 (Wenckebach) second-degree block, conduction in the atrioventricular bundle becomes progressively more impaired from beat to beat, as shown by an increasing P-R interval in the electrocardiogram, until conduction fails completely, and a ventricular beat is missed (Fig. P42). Thus, in the radial pulse, every third, fourth or fifth beat may be dropped; there is also a slight progressive increase in ventricular rate during the runs of conducted beats, but this cannot be detected by palpation alone. This type of atrioventricular block is relatively benign and may be a transient occurrence in myocardial infarction or digoxin intoxication. Normal conduction is almost always restored by exercise, atropine or any other measure that increases the atrial rate (Fig. P42). Mobitz type II second-degree block is a much more serious condition. In its mildest form, beats may be dropped intermittently, as in type I, but without any previous lengthening of the P-R interval; later, 2:1 or 3:1 block with a slow regular ventricular rhythm is common. Increasing the atrial rate, by any means, increases the severity of the block (see also PULSE, ABNORMAL RATE, p. 537).

Two mechanical causes of a palpably irregular pulse should be mentioned. Pulsus alternans is characterized by a regular rhythm with alternation in amplitude of the equally spaced beats. It may be easily palpable but is more often detected by sphygmomanometry. As the cuff pressure is lowered, half of the beats are heard at the higher systolic pressure and, as the pressure of the smaller amplitude beats is reached, the rate seems suddenly to double. A run of pulsus alternans is often initiated by an extrasystole, but it must not be confused with pulsus bigeminus. Nor should it be confused with electrical alternans in which the QRS complexes alternate in amplitude; the two conditions may co-exist, however. Although rather rare, pulsus alternans is an important sign of left ventricular failure.

Pulsus paradoxus is a feature of constrictive pericarditis, high-pressure pericardial effusion and other less common conditions in which the primary abnormality is failure of the ventricles to fill adequately (restrictive cardiomyopathy). It is also important evidence of the severity of an attack of asthma. The normal tendency of the pulse pressure to fall slightly during inspiration is much exaggerated so that, in a gross example, the arterial pulse may become impalpable during inspiration. At first, the impression
is of a grossly irregular pulse, but there is little difficulty in relating the changes to the phases of respiration; the diagnosis is confirmed by finding other evidence of pericardial constriction. This includes a rise in the already elevated venous pressure on inspiration instead of the usual fall, known also as Kussmaul's sign. The arterial changes are an exaggeration of, not opposite to, the normal; the term 'paradoxical' was applied because the heart's action appeared to have ceased during inspiration and yet, paradoxically, heart sounds could still be heard normally at this time.

PULSES, UNEQUAL

Gerry Carr-White

A thorough physical examination should include palpation of all the easily accessible arterial pulses. The pulses on the two sides should be compared and, in the arms, inequalities should be confirmed by sphygmomanometry. It must be remembered that, in normal subjects, the blood pressure in the right arm may be slightly higher than that in the left; this difference is, however, rarely palpable. It is always worth recording the arteries in which the pulse has been felt, if only for future reference; the significance of an absent pulse is much greater if it is known to have been present on a previous occasion.

The pulse in one or other radial artery may be reduced or absent as a result of a minor congenital abnormality in the course or calibre of the vessel. Other congenital conditions in which the radial pulses may be unequal include a few cases of coarctation of the aorta. In 2 per cent of cases, the lesion is proximal to the left subclavian artery, so that the pulses in the left arm are weaker than those in the right; in addition, stenosis of the origin of a subclavian artery is a rare complication of coarctation.

Inequality of pulses previously known to be equal is a most important sign. In the legs, atherosclerosis of the larger arteries is the most common cause, and the level of occlusion should be sought by comparing the pulses in the femoral, popliteal, posterior tibial and dorsalis pedis arteries. Another cause of unequal pulses, usually in the legs, is Buerger's disease, also known as thromboangiitis obliterans. Atherosclerosis is less common in the vessels of the upper limbs, but it can certainly involve the branches of the aortic arch and cause inequality of the brachial and radial pulses. Giant-cell arteritis and other inflammatory diseases of arteries occasionally cause occlusion of major limb vessels.

Arterial embolism is an important cause of unequal pulses in the upper or lower limbs. The three most
common sources of such an embolism are the left atrium in atrial fibrillation, particularly in association with mitral valve disease; vegetations in infective endocarditis of the mitral or aortic valve; and mural thrombus laid down on the endocardial surface of a myocardial infarct. Less common causes of systemic embolism are left atrial myxoma, left ventricular endocardial thrombus in a ventricular aneurysm or in dilated cardiomyopathy, thrombus detached from an atherosclerotic plaque in the aorta, and a so-called ‘paradoxical embolus’ passing from veins in the legs via a patent foramen ovale to the systemic circulation. This is more likely to occur if the pressures on the right side of the heart have been raised, often by a previous pulmonary embolism.

Arterial thrombosis of major vessels is a less common but well-recognized cause of unequal pulses. Its most frequent cause is thrombus due to aortic aneurysm, particularly of the thoracic root, which may manifest with acute limb ischaemia, giving rise to unequal pulses. Pseudoxanthoma elasticum is a rare cause of occlusive arteriopathy involving mainly the coronary and peripheral circulations and leading to unequal lower limb pulses. Frequent palpation of the arterial pulses is most important when dissecting aneurysm of the aorta is suspected. As the dissection proceeds along the length of the aorta, the branches may be occluded one by one over a period of a few hours; the process may be capricious, branches past which the dissection has spread unexpectedly remaining patent. If re-entry occurs, the pulse may return in arteries previously occluded.

Takayasu’s disease, or the ‘pulseless disease’, is a rare form of arteritis involving the branches of the aortic arch. Apart from inequality in the pulses in the arms or, perhaps more commonly, obliteration of the pulses in both arms, signs and symptoms of cerebral ischaemia are common. Takayasu’s disease is one of the causes of so-called ‘reversed coarctation’, a not very satisfactory term implying diminished or absent pulses in the arms with normal femoral pulses. This situation can also occur as a result of aortic aneurysm, particularly of the arch, in which unequal pulses in the arms may be of diagnostic importance.

Occlusion of a subclavian artery by external pressure, such as by a cervical rib or a tumour in that region, must be remembered. In cervical rib particularly, pain, paraesthesiae and weakness and wasting of the small muscles of the hand, due to compression of the first thoracic root, are common associated features.

Iatrogenic causes of unequal pulses include intraarterial cannulation and drug administration, previous subclavian–pulmonary artery anastomosis for cyanotic congenital heart disease, and brachial arteriotomy.

PUPILS, ABNORMALITIES OF

David Werring & Mark Kinirons

Abnormalities of the pupil are a vital aid to neurological and ophthalmological diagnosis. Careful examination of the shape, size and reactions of the pupils provides important information about the integrity of the autonomic (parasympathetic and sympathetic) pathways, the anterior visual pathways and the brainstem. No special equipment is needed to examine the pupils – simply a bright torch, a dark environment, and a few minutes to allow accurate observations to be made. It is important to have an understanding of normal pupil function.

ANATOMY

Normal pupil function depends on the integrity of the pupillary light reflex pathway. This pathway comprises afferent and efferent limbs. The afferent limb consists of retinal receptors; ganglion cell axons in the optic nerve; the optic chiasma; the optic tract (excluding the lateral geniculate body); and the pretectal nucleus of the midbrain. From the pretectal nucleus, interneurones stimulate the pupilloconstrictor motor (Edinger–Westphal) nuclei on both sides of the midbrain. The efferent limb begins with parasympathetic fibres that pass in the IIIrd (oculomotor) cranial nerve to the ciliary ganglion and then the short ciliary nerves. This system is affected by the amount of light falling on the retina, the degree of retinal light adaptation, the accommodation effort of the eyes, and input from the frontal and occipital cortex and the reticular formation. A lesion in one afferent limb diminishes the input to the ipsilateral pretectal nucleus, which is relayed to both Edinger–Westphal nuclei. Thus, a lesion in one afferent limb (e.g. the optic nerve) decreases pupillary constriction in both eyes when the affected side is stimulated compared with when the normal side limb is stimulated. The ipsilateral response is termed the ‘direct pupillary reaction’; the contralateral response is termed the ‘consensual pupillary reaction’. An afferent lesion can be detected clinically during the swinging light test as a relative afferent pupillary defect (RAPD). A bright light is shone into the affected eye and maintained until the
pupillary constriction is static, and then rapidly into the unaffected eye, with a similar reaction. When the light is swung again rapidly from the normal to the affected eye, there is a paradoxical dilatation of the pupil (Marcus Gunn phenomenon). In clinical practice, a RAPD is diagnostic of an incomplete optic nerve lesion. Pupil sizes are always equal as long as the output signals (afferent limb) are equal, so an isolated afferent pathway lesion does not give papillary size inequality (anisocoria).

REGULATION OF PUPIL SIZE
The size of the pupil is determined by the balance between two opposing iris muscles: the circular sphincter (pupilloconstrictor) muscle (under parasympathetic control) and the longitudinal radial pupillodilator muscle (under sympathetic control). As described above, the two-neurone parasympathetic pathway begins in the Edinger–Westphal (pupillomotor constrictor) subnucleus of the oculomotor complex, travelling to the ciliary ganglion, where it synapses with the post-ganglionic short ciliary nerves innervating the iris pupilloconstrictor muscle. The sympathetic pathway consists of a three-neurone chain, originating in the posterolateral hypothalamus. The first neurone traverses the lateral brainstem, descending to the spinal cord segments C8–T2. The second neurone ascends across the lung apex and then the neck to synapse in the superior cervical ganglion. The third (post-ganglionic) neurone follows alongside the internal carotid artery into the skull (carotid canal) and exits the superior orbital fissure with the ophthalmic division of the trigeminal (Vth) nerve. The first neurone travels to synapse in the superior cervical ganglion. The second neurone ascends across the lung apex and then the neck to synapse in the superior cervical ganglion. The third (post-ganglionic) neurone follows alongside the internal carotid artery into the skull (carotid canal) and exits the superior orbital fissure with the ophthalmic division of the trigeminal (Vth) nerve. The sympathetic pathway continues to the nasociliary branch to the pupilloconstrictor iris muscle.

Differences in pupil size are common: about 20 per cent of the normal population have a physiological anisocoria.

ABNORMALITIES OF PUPIL SHAPE
The normal pupil is circular or slightly oval, with its longer axis horizontal. An irregularly shaped pupil is due to local damage to the iris. The circular shape of the pupil is also lost in coloboma. Common causes include adhesions between the iris and the lens due to previous iritis, trauma or surgery. If an iris anomaly or iris damage is suspected from the patient’s history or examination (iris atrophy, distorted pupillary margin or heterochromia), the patient is most appropriately referred to an ophthalmologist. Anisocoria in eyes with normal iris muscles can be presumed to have pathological damage to the nerves innervating the iris muscles – that is, anisocoria of neurological origin. This is subdivided into two groups: (i) anisocoria that is greater in darkness (when the abnormal pupil is smaller because of abnormal dilatation); and (ii) anisocoria that is greater in light (when the abnormal pupil is the larger because of abnormal constriction).

ABNORMALITIES OF PUPIL SIZE
Enlargement of the pupil is termed mydriasis; constriction of the pupil is termed miosis.

Abnormally small pupil (anisocoria greater in darkness)
Physiological anisocoria
Anisocoria, usually of less than 1 mm, occurs in about 20 per cent of the general population and may be of no pathological significance. The anisocoria may alternate between sides. More pronounced difference in the size of the pupils is likely to be symptomatic of an organic lesion.

Horner’s syndrome
This is a defect in the sympathetic innervation of the eye (see Fig. P.26). The full syndrome consists of ptosis, miosis and anhydrosis. The upper lid ptosis is mild, and lower lid ptosis creates the false impression of enophthalmos. The anisocoria is of the order of 1 mm, and is less in bright light and greater in darkness. The most specific clinical sign of Horner’s syndrome is dilatation lag of the meiotic pupil compared with the normal pupil when viewed over 15–20 seconds in darkness. The pupillary light reflex and the accommodation reflex are preserved.

Aberrant regeneration
After the oculomotor nerve suffers a traumatic or compressive injury, the regenerating axons may grow along an aberrant course. Axons originally destined to supply extraocular muscles may instead sprout to innervate the iris sphincter. Therefore, whenever the patient uses the extraocular muscle concerned, the pupil constricts – a type of synkinesis. Any of the oculomotor nerve branches that innervate extraocular muscles can aberrantly innervate the iris sphincter.

Old Adie’s (tonic) pupil
A tonic pupil (see below) that has been denervated for years eventually becomes smaller than the normal pupil and does not dilate properly. This reason for this is not known.

Pontine stroke
In pontine stroke there is bilateral pupillary constriction. Opiate drugs can cause a similar papillary appearance.
**PUPILS, ABNORMALITIES OF**

**Abnormally large pupil (anisocoria greater in bright light)**

*Adie’s (tonic) pupil*

This is due to injury of the ciliary ganglion or short ciliary nerves, and post-ganglionic parasympathetic denervation of the iris sphincter and ciliary muscles. Because the iris sphincter constricts the pupil and the ciliary muscle regulates accommodation (near vision), the affected pupil is large and reacts poorly to light and accommodation, so that the patient develops difficulty with close work. After several weeks, the injured post-ganglionic parasympathetic fibres (short ciliary nerves) begin to sprout collaterals and regenerate. Because there are more accommodation fibres than pupilloconstrictor fibres, these predominate. Some accommodation fibres are imperfectly directed to the sphincter muscle, resulting in segmental areas of palsy and constriction of the iris sphincter with *vermiform movements* and a slow (tonic) contraction of the sphincter whenever the patient attempts to accommodate. The iris remains poorly responsive to light, so there is light–near dissociation. Over months and years, the pupil becomes smaller.

**Oculomotor (Illrd) nerve palsy**

The oculomotor nerve supplies the iris sphincter and ciliary muscles, levator palpebrae, superior rectus, inferior rectus, medial rectus and inferior oblique extraocular muscles. The clinical features of oculomotor palsy are therefore ptosis, pupil dilatation (mydriasis) and ophthalmoplegia. The patient will be able to abduct the affected eye if the VIth (abducens) nerve is intact but will have impaired movement in all other directions. The unopposed lateral rectus action will cause the eye to be deviated outward (exotropia) at rest. The upper lid ptosis can be mild or complete. A compressive lesion (‘surgical third nerve palsy’) is likely to affect the fibres to the sphincter muscle since these traverse the outer aspect of the oculomotor nerve. The pupil is larger than the normal pupil and reacts poorly to light and accommodation. The most common cause of a pupil-involving Illrd nerve palsy is a posterior communicating artery aneurysm, in which the pupil is involved in about 80–90 per cent of cases. In the comatose patient in whom assessment of lid and eye movement function is difficult, pupillary enlargement may be the most important sign of tentorial herniation (‘coning’). If the pupillary size and responses are spared (‘non-surgical third nerve palsy’), a process affecting the inner oculomotor fibres is more likely. The most common cause of a pupil-sparing Illrd nerve palsy is a ‘microvascular’ lesion, often in the context of diabetes. Other causes include mononeuritis due to vasculitis, granuloma (e.g. sarcoidosis) and treponemal infection. It should be remembered, however, that up to 40 per cent of microvascular oculomotor palsies may show pupillary involvement. A fixed dilated pupil that remains an isolated abnormality for more than 1 week in a neurologically intact patient is probably not an acute oculomotor nerve palsy.

**Pharmacological mydriasis**

Topical agents that can pharmacologically dilate the pupil fall into two classes: (i) sympathomimetics, which stimulate the pupillodilator; or (ii) anticholinergics, which inhibit the pupilloconstrictor. Topical agents include phenylephrine, cocaine, hydroxyamphetamine, guanethidine, atropine, scopolamine, cyclopentolate and tropicamide. Inadvertent exposure to mydriatic agents may occur from eye drops used for red eyes, scopolamine skin patches used for motion sickness, anticholinergic agents used for asthma, or certain plants. Occasionally, mydriatics may be deliberately instilled by patients, particularly those in medical or allied professions, in order to mimic true pathology.

**An isolated fixed, dilated pupil**

This clinical finding deserves special mention as it sometimes causes unnecessary emergency assessment of a patient. Confirmation that the dilated pupil is an isolated clinical finding is important – there are truly no other signs of focal neurological deficit, ipsilateral ptosis or ophthalmoplegia. If the patient complains of diplopia, it should be assumed that there is weakness of one or more extraocular muscles. An early oculomotor palsy may have very subtle ocular motility abnormalities. If no ophthalmoplegia is present, the two most common causes of an isolated, fixed dilated pupil are tonic pupil or pharmacologically induced mydriasis. Rare causes include a fascicular or very early compressive Illrd nerve palsy. If there is better accommodation than light response (light–near dissociation), the likely diagnosis is an Adie’s (tonic) pupil. If the dilated pupil is unreactive to light or accommodation, an acute tonic pupil is still possible, but pharmacological mydriasis must be considered. Pilocarpine (1%) will usually constrict a tonic pupil or oculomotor palsy, but not a pharmacologically manipulated pupil.

**Transient pupillary abnormalities**

The pupil varies in size depending on age. In infancy it is small, but it becomes larger during young adult and middle life, and then small again in old age. As a general rule, the pupil is smaller in hypermetropic (long-sighted) and larger in myopic (short-sighted) eyes.
Ophthalmoplegic migraine occurs in children and adolescents, and typical migraine is followed by a unilateral oculomotor nerve palsy. Ophthalmoplegia and pupillary involvement are usual, and the deficit takes 1–4 weeks to resolve. The diagnosis is based on a characteristic history and negative evaluation for other possible causes of an oculomotor palsy.

Benign episodic unilateral mydriasis has been noted in young women with migraines, and it lasts for an average of 12 hours; it recurs with a frequency of two to three attacks per month.

Tadpole-shaped pupils due to intermittent segmental spasm of the dilator muscle last for a few minutes and recur several times daily or weekly. The episodic distortion eventually resolves spontaneously. Conditions associated with tadpole pupils are Horner’s syndrome, tonic pupil and migraines.

Hippus is a benign cause of spontaneous papillary movement; both pupils are seen to constrict and dilate simultaneously, without any obvious stimulus being applied. This is simply an exaggeration of physiological variation in pupil size due to fluctuations in the different inputs to the pupillomotor pathways.

**PURPURA**

Mark Kinirons

(See also BLEEDING, p. 56; BRUISES, p. 76.)

The presence of purpura usually indicates thrombocytopenia (Box P 14), although there are, additionally, non-thrombocytopenic causes (Box P 15). In any patient with purpura due to a reduced platelet count, it is important to ascertain the cause and severity because purpura usually indicates a severe haemorrhagic potential. Whereas purpura on the skin is not life-threatening, thrombocytopenia can result in rapidly fatal intracranial or massive gastrointestinal haemorrhage. Purpura usually indicates a platelet count of less than 30 × 10^9/l.

Platelet functional disorders (e.g. thrombasthenia or platelet storage pool disorder) do not usually result in purpura unless there is a secondary provocative stimulus (e.g. protracted coughing). The presence of purpura in an individual with a normal platelet count is indicative of a vasculitis (Fig. P 43). Whereas thrombocytopenia results in flat purpura lesions, those due to a vasculitis are often slightly raised, and on some occasions may be up to 1 cm in diameter.

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**Box P 14** Causes of thrombocytopenia

<table>
<thead>
<tr>
<th>Impaired production of platelets</th>
<th>• Drugs causing immune destruction</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Congenital megakaryocytic abnormalities</td>
<td>• Quinine</td>
</tr>
<tr>
<td>• Wiskott–Aldrich</td>
<td>• Gold</td>
</tr>
<tr>
<td>• May–Hegglin anomaly</td>
<td>• Penicillins</td>
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<tr>
<td>• Thrombocytopenia with absent radii</td>
<td>• Para-aminosalicylic acid</td>
</tr>
<tr>
<td>• Marrow hypoplasia</td>
<td>• Rifampicin</td>
</tr>
<tr>
<td>• Aplastic anaemia</td>
<td>• Methyldopa</td>
</tr>
<tr>
<td>• Megaloblastosis</td>
<td>• Heparin</td>
</tr>
<tr>
<td>• Vitamin B12/folate deficiency</td>
<td>• Neonatal thrombocytopenia</td>
</tr>
<tr>
<td>• Folate antagonists</td>
<td>• Maternal autoantibody</td>
</tr>
<tr>
<td>• Toxins</td>
<td>• Maternal isoantibody</td>
</tr>
<tr>
<td>• Chemotherapy</td>
<td>• Post-transfusional purpura</td>
</tr>
<tr>
<td>• Drugs</td>
<td>• Disseminated intravascular coagulation</td>
</tr>
<tr>
<td>• Phenylbutazone</td>
<td>• Pregnancy-associated conditions</td>
</tr>
<tr>
<td>• Sulphonamides</td>
<td>• Eclampsia</td>
</tr>
<tr>
<td>• Chloramphenicol</td>
<td>• Abruptio placentae</td>
</tr>
<tr>
<td>• Procainamide</td>
<td>• Retained dead fetus</td>
</tr>
<tr>
<td>• Thiazides</td>
<td>• Septicaemia</td>
</tr>
<tr>
<td>• Oestrogens</td>
<td>• Hypothermia</td>
</tr>
<tr>
<td>• Ethanol</td>
<td>• Asphyxia</td>
</tr>
<tr>
<td>• Ionizing radiation</td>
<td>• Cardiopulmonary arrest</td>
</tr>
<tr>
<td>• Infiltiration by:</td>
<td>• Local thrombosis</td>
</tr>
<tr>
<td>• Leukaemia</td>
<td>• Massive thromboembolism</td>
</tr>
<tr>
<td>• Lymphoma</td>
<td>• Giant haemangiomas</td>
</tr>
<tr>
<td>• Myeloma</td>
<td>• Thrombotic thrombocytopenic purpura (Moscovitz syndrome)</td>
</tr>
<tr>
<td>• Carcinoma</td>
<td>• Haemolytic–uraemic syndrome</td>
</tr>
<tr>
<td>• Paroxysmal nocturnal haemoglobinuria</td>
<td>• Virus</td>
</tr>
<tr>
<td>• Myelodysplasia</td>
<td>• Rubella</td>
</tr>
</tbody>
</table>

**Box P 15** Non-thrombocytopenic causes of purpura

<table>
<thead>
<tr>
<th>Vasculitis</th>
<th>• Orthostatic</th>
</tr>
</thead>
<tbody>
<tr>
<td>Henoch–Schönlein purpura</td>
<td>• Senile</td>
</tr>
<tr>
<td>Systemic lupus erythematosus</td>
<td>Steroid-induced</td>
</tr>
<tr>
<td>Drugs</td>
<td>Scurvy</td>
</tr>
<tr>
<td>Mechanical</td>
<td>Factitious purpurae</td>
</tr>
<tr>
<td>Coughing</td>
<td></td>
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</tbody>
</table>
A pustule is an elevation of the skin containing pus, differing from a vesicle or bulla only in its contents, a vesicle or bulla containing clear fluid.

Pustules may develop from vesicles that have become purulent, or from papules. They may develop so rapidly that their origin cannot be observed. They vary in colour from bright yellow to cream, orange, grey or green. Examination with a hand lens may show them to arise in hair follicles, but they may occur on normal skin or around sweat pores (Box P.16). Establishing the diagnosis of a pustular lesion or rash may be assisted by determining whether or not the pus is sterile on bacteriological examination.

**PUSTULES OF INFECTIOUS ORIGIN**
The most common bacterial cause of a pustule is infection with *Staphylococcus*. Systemic factors (e.g. diabetes) may predispose, as may the effect on the skin of heat and humidity, friction, or the use of topical corticosteroids on the skin. Recurrent infection may result from chronic staphylococcal carriage in the anterior nares or the perineum. Severe recurrent lesions may occur in individuals who are carriers of the Panton–Valentine leucocidin strain of *Staphylococcus aureus*. They characteristically develop multiple deep necrotic boils, and these will not be cured until total skin decontamination has taken place. A boil beginning in the hair follicles of the eyelid is known as a stye (hordeolum).

A carbuncle resembles a boil, but the infection spreads to the deeper tissues and, when rupture occurs, there may be several openings onto the skin. Boils may follow one another in series for many months, when the condition is known as chronic furunculosis. Sycosis barbae is staphylococcal folliculitis of the hairy areas of the face, including the eyebrows. It differs in no way from the other forms of staphylococcal folliculitis; it may be localized to small areas, or it may affect the whole of the face and neck. The pustules are grouped on a bright erythematous base, and in the centre of each pustule there is a hair that pulls out easily. It is now much less common than when men were shaved in barbers’ shops and infection was transmitted from customer to customer on the razor. It must be distinguished from the more common seborrhoeic sycosis in which the predominating lesions are red papules with little or no pustule formation. In all forms of staphylococcal folliculitis, the diagnosis is fairly simple. Only rarely do other organisms (e.g. *Streptococcus* or Gram-negative organisms) produce similar lesions.

An ecthyma begins as a burning painful vesicopustule on an erythematous base, either singly or as sparse pustules, undergoes rapid necrosis, and a yellow pustule is formed, which becomes boggy, fluctuates and ultimately ruptures and discharges pus. In many cases, a hard ‘core’ of necrotic material is extruded before healing takes place. In patients with recurrent boils, microbiological investigation is mandatory. Some affected subjects may be shown to be carriers of the Panton–Valentine leucocidin strain of *Staphylococcus aureus*.
lesions scattered over the buttocks and lower legs. Later, deep-crusted ulcers form, which heal slowly with considerable scarring. Patients are usually young women, debilitated by anaemia or systemic illness or suffering from HIV disease. Streptococci or staphylococci may be cultured. Boils in the axillae and groins can be the presentation of hidradenitis suppurativa, an intractable purulent condition affecting the apocrine sweat glands. Microbiology swabs are commonly negative, and this assists in distinguishing the condition from recurrent boils.

Recurrent purulent skin infections in a child, especially with crops of pustules in the webs of the fingers and on the wrists, are sometimes a feature of neglected scabies. Pustules are also a feature of secondarily infected dermatitis and can be particularly widespread in atopic dermatitis.

Satellite vesicopustules at the edge of a moist eroded patch are highly suggestive of candidiasis (thrush). This particularly favours damp, intertriginous areas, such as the submammary, perineal and perioral folds of skin. Scrapings from lesions warmed with 10 per cent potassium hydroxide show small, oval, budding, thin-walled spores on microscopy; culture shows Candida albicans.

Some dermatophyte species appear to proliferate deep in hair follicles, especially Trichophyton mentagrophytes. Inappropriate treatment of superficial fungal infections with topical corticosteroids often leads to an apparent initial improvement of the condition, but fungal growth persists deep in the hair follicles and a curious resistant pustular eruption may result – the so-called tinea incognito. Scalp ringworm in cases where there is a vigorous inflammatory reaction may present with a discharging pustular mass (kerion). Fungal elements will be seen on microscopy. Characteristic discrete inflammatory pustules (which may become haemorrhagic) accompany the fever, arthritis and tenosynovitis of gonococcal septicaemia.

Recognition of this skin lesion can lead to making this important diagnosis at an earlier stage, particularly in women who may have little in the way of genital symptoms.

Anthrax infection of the skin in its localized variety takes the form of a carbuncle-like inflammatory lesion caused by Bacillus anthracis. It is contracted from the hides or hairs of cattle or goats, or rarely from shaving brushes. It attacks an exposed area of skin, and the incubation period is 1–3 days. The first sign is a small, itching red macule, not unlike a flea bite. In 2 days, a papule forms, which rapidly becomes a pustule; this soon ruptures, and sometimes blood as well as pus is extruded. There results a gangrenous ulcer that, in a simple case, heals in a few weeks. However, in severe cases, there may be grave constitutional symptoms with septicaemia and rapid death, and sometimes there are multiple skin lesions. It must be distinguished from a carbuncle and extragenital syphilitic chancre. Scrapings from the lesion contain the causal organism in large numbers. Anthrax is mostly an occupational disease in handlers of hides or wool, but it is endemic in some parts of the world.

Unilateral pustules near the lips may have developed from herpes simplex vesicles and tend to be greyish in colour. The grouped distribution and a history of recurrent lesions or recent respiratory tract infection are suggestive.

The pustule was once an important manifestation of smallpox. After an incubation period of 5–12 days, the disease began with fever, headache, backache and vomiting. On the third and fourth days, there was macular erythema, and after a few hours shotty papules developed. These became vesicles, and by the fifth day pustulated. The rash was most profuse on the head and limbs, and there was only one crop of lesions, which matured in ‘majestic’ concert. The vesicles were tough, firm and often multilocular, and showed definite umbilication. By the time the pustules formed, the umbilication was less well marked. In severe cases, there was confluence of the pustules, particularly on the face. The mucous membranes were often also involved. As a rule, the temperature fell slightly with the eruption but rose again on the eighth or the ninth day when the pustules ruptured. In the final stage, the pustules dried up to form brown crusts. Pitting or scarring was the rule. Previous vaccination modified the course of the disease considerably, and there was a type of the disease known as alastrim (variola minor).

Chickenpox can be distinguished by the usually milder nature of disease, which is vesicular rather than purulent, and its profusion on the trunk rather than the extremities. In chickenpox, the eruption comes out in successive crops, and the vesicles are unilocular, fragile and do not exhibit umbilication. In spite of these differences, mild smallpox and severe chickenpox were difficult to differentiate.

Other viral diseases causing pustules are vaccinia, cowpox and orf. In the so-called pustular syphilitides, there are no true pustules, their resemblance to pustules being only superficial; on incision, they will be found to be solid and contain no pus. The histology is diagnostic, and the serology will be positive.
Perhaps the most modern form of folliculitis is a new epidemic of *Pseudomonas folliculitis* that has been described in those indulging in jacuzzi-bathing. The organism is inoculated on to the skin by the high-pressure water jets.

**STERILE PUSTULES**

Pustules do not always indicate cutaneous infection and can arise during inflammatory dermatoses. These are often referred to as ‘sterile’ pustules, as routine bacteriological examination of the pus always yields negative results. The classical example is *pustular psoriasis of the palms and soles*, in which bright yellow sterile pustules arise within well-demarcated areas on palms and soles (Fig. P.44). As they age, the pustules change to dark-brown macules and eventually peel off. The condition is most common in middle-aged women, and is notoriously resistant to conventional treatment. Only 25 per cent of such patients have evidence of psoriasis elsewhere on the body. On very rare occasions, psoriasis may produce widespread sheets of sterile yellow pustules, associated with considerable toxicity and fever – a phenomenon known as generalized pustular psoriasis of von Zumbusch. The incidence of this rare and dangerous complication of psoriasis rose dramatically following the introduction of potent topical steroids; it has become uncommon again, now that steroids are used more cautiously and appropriately in treatment.

Sterile pustules on the face are seen in *acne*, *rosacea* and *perioral dermatitis*. In *acne* these are associated with evidence of comedones, acne papules and perhaps acne cysts (Fig. P.45). The distribution is more peripheral on the face than in *rosacea*, where pustules surround the ‘muzzle’ area and are associated with vascular changes (erythema, flushing and telangiectasia), as well as papules and sometimes hypertrophied and patulous pilosebaceous pores on the nose (rhinophyma). *Perioral dermatitis* is a modern condition that was unknown before the introduction of fluorinated topical corticosteroids; the pustules are tiny and surmount painful small red papules, which abound around the mouth. A further diagnostic feature is a background of perioral erythema with a halo of pallor around the lip margins.

If pustules are extremely pruritic, consideration must be given to infestation or *dermatitis herpetiformis*. Scabies has already been mentioned; the burrows can become pustular, but in *swimmer’s itch*, 12 hours after freshwater bathing, itchy weals may appear on the legs, and later vesicopustules appear. Scrapes from these may reveal the causative bird or mammal schistosomes that have been inoculated. In *dermatitis herpetiformis*, the greyish vesicopustules are transient because of rapid deep excoriation. They occur on the forehead and scalp, shoulders, buttocks, natal cleft and knees. Diagnosis is often delayed.

Widespread pustulosis over the chest and back may follow a febrile illness or travel to a tropical environment (*miliaria*). A similar monomorphic pustulation of the face and upper trunk is prone to develop 4–8 weeks after the administration of systemic steroids or adreno-corticotropic hormone, or more rapidly after iodides or bromides. A widespread sterile pustulosis of the skin is a recurrent feature of...
Behçet’s disease, in which deep, painful ulcers develop on the orogenital mucosae. A dermatologist's finest hour comes with the diagnosis of the extremely rare subcorneal pustular dermatosis of Sneddon and Wilkinson. Here, the pustules are very superficial and often form rings or gyrate patterns in flexures or on the flexor aspects of extremities. The condition affects middle-aged women, and the cause is unknown. Again, the pustules are sterile.

PYREXIA, PROLONGED

Alex West

Fever is a controlled elevation of body temperature brought about by thermoregulatory reflexes. This distinguishes fever from hyperthermia, which is an uncontrolled rise in temperature as a consequence of thermal overload, as in heat stroke or the loss of thermoregulatory function. Malignant hyperthermia is a rare, genetically determined disorder of muscle metabolism that causes a very high temperature after general anaesthesia. Certain cytokines (e.g. interleukin 1 [IL-1], IL-6 and tumour necrosis factor) produced by inflammatory cells act as endogenous pyrogenic mediators. There is evidence that the fever response may be beneficial in cases of infection, and fever is a cardinal sign of inflammation, whatever the cause may be.

A fever may be described as prolonged if it persists beyond 3 weeks. If no cause has been found after these 3 weeks with investigation, it may be referred to as a ‘pyrexia of unknown origin’ (PUO). In general it is worth remembering that the three main causes when ultimately discovered fall relatively evenly between infections, malignancies and inflammatory disorders (with some regional variation depending on country). It may be persistent, remittent, swinging up and down, but remaining above normal; or intermittent, with periods of normal temperature between febrile episodes. In many cases, the cause may be obvious, but in others further investigation is required, for example:

- **Blood counts:** a high polymorphonuclear leucocytosis is not invariably present with local abscess formation, but a full blood count may help to explain many fevers
- **Serum agglutination tests**, for example for typhoid fever, paratyphoid fever, abortus fever, listeriosis, etc.
- **Blood cultures**
- **Urine cultures**
- **Bacterial examination of exudates**
- **Lumbar puncture**, revealing, for example, unexpected meningeal infection
- **X-ray examination of the thorax, kidney, colon and elsewhere, to identify or exclude infection or neoplasm**
- **Computed tomography, bronchoscopy, sigmoidoscopy or endoscopy**
- **Blood and tissue cultures**, from bone, spleen, liver or other tissues, for infective organisms, and biopsy for evidence of malignancy or specific inflammatory tissue changes

CAUSES OF PROLONGED PYREXIA

The list of conditions that may cause prolonged pyrexia is lengthy. The relative frequency of the different causes varies greatly from country to country and, to a lesser extent, from time to time. The more common conditions to consider are listed in Box P.17. Inevitably, such a list is bound to be incomplete and cannot be comprehensive. Space does not permit a full description of each of the diseases mentioned, but the following are some of the salient points.

**Box P.17 Causes of prolonged pyrexia**

<table>
<thead>
<tr>
<th>Specific fevers</th>
<th>Localized infection</th>
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<tbody>
<tr>
<td>Viral</td>
<td>Prostatic abscess</td>
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<tr>
<td>Salmonella</td>
<td>Ischorectal abscess</td>
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<tr>
<td>S. typhi</td>
<td>Pyosalpinx</td>
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<tr>
<td>S. paratyphi</td>
<td>Suppurating ovarian cyst</td>
</tr>
<tr>
<td>Rickettsia</td>
<td>Parametritic abscess</td>
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<tr>
<td>Rocky Mountain spotted fever</td>
<td>Empyema of the maxillary antrum, frontal sinus or ethmoidal air cells</td>
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<tr>
<td>Q fever</td>
<td>Osteomyelitis</td>
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<tr>
<td>Trench fever</td>
<td>Infected lymph nodes in neck, axilla or groin</td>
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<tr>
<td>Legionella</td>
<td>Mammary or submammary abscess</td>
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<tr>
<td>Legioniare’s disease</td>
<td>Empyema thoracis</td>
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<tr>
<td>Chlamydia</td>
<td>Lung abscess</td>
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<td>Psittacosis</td>
<td>Hepatic abscess</td>
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<td>Fungus</td>
<td>Renal abscess</td>
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<tr>
<td>Coccidioidomycosis</td>
<td>Splenic abscess</td>
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<tr>
<td>Histoplasmosis</td>
<td>Empyema of gallbladder</td>
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<tr>
<td>Spirochaetes</td>
<td>Suppurative cholangitis</td>
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<tr>
<td>Syphilis</td>
<td>Suppurative pyelonephritis</td>
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<tr>
<td>Leptospirosis</td>
<td>Subdiaphragmatic abscess</td>
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<tr>
<td>Lyme disease</td>
<td>Bronchiectasis</td>
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<tr>
<td>Brucella</td>
<td>Appendix abscess</td>
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<tr>
<td>Abortus and melitensis fever</td>
<td>Perihepatic abscess</td>
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<tr>
<td>Tuberculosis</td>
<td>Diverticulitis</td>
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<tr>
<td>Listeriosis</td>
<td>Lumbar and iliac retroperitoneal abscess</td>
</tr>
<tr>
<td>Streptobacillary rat-bite fever</td>
<td>Psosas abscess</td>
</tr>
<tr>
<td>Tularaemia</td>
<td>Actinomycosis of jaw, cheek, neck, lung, liver, spine or caecum</td>
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(Continued)
SPECIFIC FEVERS

Today, infectious mononucleosis is one of the most common causes of prolonged fever in children and young adults, and diagnosis is often difficult. Paul–Bunnell or ‘Monos’ tests are useful in diagnosis after the first 2 weeks or so. These are tests for heterophile antibodies; several specific antibodies against components of the Epstein–Barr virus (EBV) have also been identified.

Infectious mononucleosis should be seriously considered in any prolonged, low-grade fever with malaise and loss of weight in children and young adults, as lymphadenopathy, splenomegaly and typical blood counts are not invariably present in all cases, or may have been present earlier and have disappeared. Some authors consider infectious mononucleosis to be a clinical syndrome rather than a single entity, as the same picture can be due to EBV, cytomegalovirus or Toxoplasma. However, as only EBV gives a positive Paul–Bunnell test, full serological tests should be carried out in all cases.

Typhoid fever suggests itself when a patient who was previously in good health suffers from a progressive fever of considerable and increasing degree with a pulse rate that is relatively slow in relation to the temperature. The illness starts with headache and malaise, but without any conspicuously abnormal signs. During the first week, the temperature rises each night to a slightly higher level than that of the previous night, until a maximum is attained and maintained during the second week. After this time, there is a progressive diminution during the third week until normal temperature is reached again. Diarrhoea may occur with foul-smelling stools of pea-soup consistency, but constipation is more usual. Abdominal pain is usually confined to the right lower quadrant. The headache, which is in most cases a conspicuous feature, persists for about a week, at which time it almost invariably ceases, thus contrasting with the headache of tuberculous meningitis. Blood cultures are positive during the first 10 days and in acute relapse; similar cultures later prove negative, although urine and faecal cultures may be positive. The spleen becomes palpable early in the disease and remains so until defervescence; it enlarges again in a relapse. Typical typhoid rose spots appear – chiefly on the abdomen, less often on the chest or back, and seldom on the limbs – from the seventh day onwards, and in successive crops. They are about 2–3 mm in diameter, rose-red in colour, fade on pressure and are without a central punctum. The majority of patients, but not all, develop a rise in agglutinins against the O antigens of the typhoid bacillus during the course of the disease. Another help in diagnosis is the absence of leucocytosis, and in the differential leucocyte count the lymphocytes are relatively increased, the polymorphonuclear cells being absolutely reduced. The leucocyte count may, however, be influenced by complications. Rigors are exceptional, a fact that sometimes helps in the diagnosis from conditions such as septicemia and malaria. Some difficulty arises in the diagnosis of those cases in which there has been previous immunization of the patient by anti-typhoid inoculations; the fever is then of shorter duration, and the illness relatively mild.

The rickettsial disorders are numerous, ranging from epidemic typhus to trench fever. Different Rickettsiae

Box P.17 (Continued) Causes of prolonged pyrexia

<table>
<thead>
<tr>
<th>Infective and inflammatory conditions</th>
<th>Infective and inflammatory conditions</th>
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<tbody>
<tr>
<td>Pyelonephritis and other urinary infections</td>
<td>Pyelonephritis and other urinary infections</td>
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<td>Papillary necrosis</td>
<td>Papillary necrosis</td>
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<td>Chronic cystitis</td>
<td>Chronic cystitis</td>
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<td>Chronic cholecystitis</td>
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<td>Phlebitis</td>
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<td>Thyroiditis</td>
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<td>Pneumonia and pneumonitis</td>
<td>Pneumonia and pneumonitis</td>
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<td>Bronchopneumonia</td>
<td>Bronchopneumonia</td>
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<td>Parametritis</td>
<td>Parametritis</td>
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<td>Vesiculitis</td>
<td>Vesiculitis</td>
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<td>Dysenteric colitis, bacillary or amoebic</td>
<td>Dysenteric colitis, bacillary or amoebic</td>
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<td>Ulcerative colitis</td>
<td>Ulcerative colitis</td>
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<td>Crohn’s disease (regional enteritis)</td>
<td>Crohn’s disease (regional enteritis)</td>
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<td>Pancreatitis</td>
<td>Pancreatitis</td>
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<td>Familial Mediterranean fever</td>
<td>Familial Mediterranean fever</td>
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<td>Sarcoidosis</td>
<td>Sarcoidosis</td>
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<td>Non-parasitic hepatic affections</td>
<td>Non-parasitic hepatic affections</td>
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<td>Cirrhosis</td>
<td>Cirrhosis</td>
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<td>Secondary carcinoma</td>
<td>Secondary carcinoma</td>
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<td>Hepatitis, subacute and chronic</td>
<td>Hepatitis, subacute and chronic</td>
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<td>Connective tissue disorders</td>
<td>Connective tissue disorders</td>
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<td>Rheumatoid arthritis</td>
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<td>Rheumatic fever</td>
<td>Rheumatic fever</td>
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<td>Systemic lupus erythematosus</td>
<td>Systemic lupus erythematosus</td>
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<td>Polyarteritis nodosa and other arteritides</td>
<td>Polyarteritis nodosa and other arteritides</td>
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<td>Polymyositis and dermatomyositis</td>
<td>Polymyositis and dermatomyositis</td>
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<td>Giant-cell arthritis</td>
<td>Giant-cell arthritis</td>
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<tr>
<td>Still’s disease (childhood and adult)</td>
<td>Still’s disease (childhood and adult)</td>
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<td>Blood diseases</td>
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<td>Aplastic anaemia</td>
<td>Aplastic anaemia</td>
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<td>Agranulocytosis</td>
<td>Agranulocytosis</td>
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<td>Lymphatic, myeloid or monocytic leukaemia, acute or chronic</td>
<td>Lymphatic, myeloid or monocytic leukaemia, acute or chronic</td>
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<td>Haemolytic anaemias</td>
<td>Haemolytic anaemias</td>
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<td>Diseases of the tropics and subtropics</td>
<td>Diseases of the tropics and subtropics</td>
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<tr>
<td>Trypanosomiasis</td>
<td>Trypanosomiasis</td>
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<td>Malaria</td>
<td>Malaria</td>
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<td>Kala-azar</td>
<td>Kala-azar</td>
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<td>Plague</td>
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<td>Relapsing fever</td>
<td>Relapsing fever</td>
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<td>Filariasis</td>
<td>Filariasis</td>
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<td>Leprosy</td>
<td>Leprosy</td>
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<td>Schistosomiasis</td>
<td>Schistosomiasis</td>
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<tr>
<td>Meningeal and cerebral haemorrhage</td>
<td>Meningeal and cerebral haemorrhage</td>
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<td>Skin conditions</td>
<td>Skin conditions</td>
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<tr>
<td>Pemphigus</td>
<td>Pemphigus</td>
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<tr>
<td>Severe or exfoliative dermatitis</td>
<td>Severe or exfoliative dermatitis</td>
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<td>Bullous pemphigoid</td>
<td>Bullous pemphigoid</td>
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<tr>
<td>Malignancy</td>
<td>Malignancy</td>
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<td>Lymphoma (e.g. Hodgkin’s disease, lymphosarcoma)</td>
<td>Lymphoma (e.g. Hodgkin’s disease, lymphosarcoma)</td>
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<td>Sarcoma</td>
<td>Sarcoma</td>
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<tr>
<td>Carcinoma</td>
<td>Carcinoma</td>
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<tr>
<td>Allergic (antigen–antibody reactive) conditions</td>
<td>Allergic (antigen–antibody reactive) conditions</td>
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<tr>
<td>Henoch–Schönlein syndrome</td>
<td>Henoch–Schönlein syndrome</td>
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<td>Allergic skin rashes</td>
<td>Allergic skin rashes</td>
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<tr>
<td>Post-cardiac injury syndrome</td>
<td>Post-cardiac injury syndrome</td>
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<tr>
<td>Factitious pyrexia produced by malingering</td>
<td>Factitious pyrexia produced by malingering</td>
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<tr>
<td>Drug reactions</td>
<td>Drug reactions</td>
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<td>Sulphonamides</td>
<td>Sulphonamides</td>
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<td>Antibiotics</td>
<td>Antibiotics</td>
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<td>Arsenicals</td>
<td>Arsenicals</td>
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<td>Iodides</td>
<td>Iodides</td>
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<tr>
<td>Barbiturates, etc.</td>
<td>Barbiturates, etc.</td>
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are transmitted by lice, fleas, ticks and mites via man, wild rodents, domestic animals and cattle. In the history of mankind, they rank high as a cause of epidemic disease causing great suffering and death, but they do not usually persist beyond 20 days and rarely cause prolonged fever. 

Legionnaires' disease usually resolves or proves fatal within 3 weeks, but the pneumonia caused may extend to the lobes of both lungs and prolong the febrile disorder. The causative organism resembles a Rickettsia in some cultural characteristics, but it is larger in size and does not react with the standard rickettsial antigens in complement fixation tests. Diagnosis via the Legionella direct antigen test performed on urine is now possible. 

Psittacosis or ornithosis is transmitted to man from parrots and the parrot family, pigeons and a number of other birds, including ducks, turkeys and chickens. It is due to a Gram-negative obligate intracellular parasite, Chlamydia psittaci. The illness may be transmitted from the patient to others by contact. Clinically, the illness is similar to typhoid fever, with liability to serious pulmonary complications. The fever lasts for about 3 weeks, and tends to end abruptly followed by a slow convalescence, but it may last for as long as 3 months. When the disease is contracted from parrots or parakeets, it tends to be more severe and prolonged. The diagnosis is suggested in a patient with an illness that bears a general resemblance to typhoid fever, but whose blood does not give the agglutination test, and especially if there has been contact with a recently imported parrot, budgerigar or pigeon. The organism may be isolated from the blood or sputum. A rising titre of complement-fixing antibody in the patient's blood is useful in diagnosis. 

Secondary syphilis may be a febrile illness. The diagnosis becomes obvious when the rash is associated with a fading primary sore, typical snail-track ulcers on the tonsils, fauces and pharynx, and generalized enlargement of most of the palpable lymph nodes. Spirochaetes may be isolated from the skin lesions. 

Lyme disease is caused by a tick-borne spirochaete, Borrelia burgdorferi. Cases have been reported from most parts of the USA, Scandinavia and Europe. The early disease is characterized by the annular skin lesion, erythema chronicum migrans (ECM). At this stage, there may be fever, lymphadenopathy, myalgia and arthralgia. A history of a tick-bite is often obtained. After several weeks of this flu-like illness, neurological, cardiac and joint changes can develop, with the latter often becoming permanent. The diagnosis can be confirmed serologically and by culture of the organisms or their demonstration in skin biopsies of ECM. 

Brucellosis is a not uncommon cause of prolonged unexplained pyrexia, usually lasting for several months and occasionally for a year. It is due to infection by Brucella abortus. This organism causes fatal abortion in cows and, apart from the geographical circumstances, the fever is identical to Mediterranean fever, in which the disease is communicated by goat's milk. Infection may arise from the ingestion of milk or the handling of infected animals or excreta. This infection may underlie obscure, long-continued febrile illness either in children or in adults. There are no characteristic symptoms, although arthritis is a frequent accompaniment. Diagnosis is usually by agglutination tests, and rarely by blood cultures. Rarely, the organism causes endocarditis, which is sometimes fatal. Cultures taken from bone marrow or liver may be positive when blood cultures are negative. Brucellar infections are particularly persistent, probably due to the intracellular location of the organism in the reticuloendothelial tissues. 

Melitensis, Mediterranean or Malta fever is one of the most prolonged of the fevers due to a known specific organism, in this case Brucella melitensis. In the undulant form of the disease, successive exacerbations of pyrexia may prolong the illness into the 16th, 18th or 20th week, or longer (Fig. P46). The condition may simulate typhoid fever, including the enlargement of the spleen and the paucity of abnormal physical signs, but there are no rose spots or other eruption. The diagnosis may be suggested by geographical factors, for example, recent residence in some parts of the Mediterranean coast or islands, or Spain, Portugal, the Canary Islands or parts of South America – and especially if the patient has been drinking goat's milk, by which the infection is transmitted. The diagnosis is established by serum-agglutination tests, which may be positive from the fifth day onwards, the patient's diluted blood serum agglutinating cultures of Brucella melitensis. Blood cultures should be carried out. Brucella suis infections, contracted from pigs, are also diagnosed by agglutination tests and blood cultures. 

Tuberculosis lesions often occur without pyrexia. On the other hand, the occurrence of some degree of fever without any apparent cause may be the sole evidence of such disease, especially the miliary form. Pulmonary tuberculosis is almost invariably pyrexial, but in an earlier stage there are often long periods of apyrexia even when the tuberculous process is active, with brief febrile spells. Tuberculous disease of the joints may be apyrexial unless secondarily infected.
Figure P.46 Temperature chart of a case of Mediterranean fever of undulant type (*Brucella melitensis*).
Glandular tuberculosis is less likely to be pyrexial when the nodes involved are cervical or bronchial than when the mesenteric and other abdominal nodes are caseous and softening (‘tabes mesenterica’), when there may be – and usually is – prolonged pyrexia. The diagnosis may be easy if there is ascites in a child or if there are palpable abdominal masses, but it may be difficult in the absence of lymph nodes and ascites in a condition of ill-health with pyrexia and with vague abdominal pains, which may be mistaken for some other non-tuberculous abdominal disease. The patient is usually young, and in European countries may be an immigrant. The presence of enlarged cervical lymph nodes supports this diagnosis.

Listeriosis is an infectious disease of animals and man worldwide. Its distribution is due to a Gram-positive bacillus, Listeria monocytogenes. Clinically, the condition may present as a meningitis, or resemble infectious mononucleosis with pharyngitis and diffuse lymphadenopathy, influenza or miliary tuberculosis. Untreated severe cases with meningitis often prove fatal. It may occur in infancy or in pregnancy. Diagnosis rests on isolating the microorganism, which resemble diphtheroid bacilli in culture, and on rising agglutination titres in the serum.

Streptobacillary rat-bite fever is caused by bites from rats and, sometimes, mice, cats, dogs and weasels. The causative organism is Streptobacillus moniliformis or Spirillum minus, and the condition is characterized by acute febrile attacks at fairly regular intervals of a few days. These persist for from 2 to 10 months.

Tularaemia is uncommon in England, but cases have occurred among those handling live rodents, for example in bacteriological laboratories. It is a specific infectious disease due to Pasteurella (Francisella) tularensis and is transmitted from rodents by ticks or deer flies, by the handling or ingestion of infected animal tissues, or by the inhalation of infected aerosols. It is not transmitted directly from human to human. A sore on a finger is commonly the start, the sore becoming a small ulcer in a day or two, with associated enlargement of the epitrochlear and axillary lymph nodes; a chill or rigor is usual, and pyrexia continues for 2–3 weeks with marked prostration followed by slow convalescence. There may be erythematous blotches on the skin, or even purpura. Swallowing a very large number of bacilli may cause a typhoid-like disorder with high fever, abdominal pain and toxicity. Lung involvement may occur, in addition to enlargement of the cervical glands. Rigors seen initially in severely ill patients may be followed by pyrexia persisting for several weeks.

The diagnosis can be confirmed by a skin test, which becomes positive in the first week, or the organism can be recovered from a mucocutaneous ulcer or regional lymph node, and occasionally from sputum on appropriate culture. Specific agglutinins appear in the serum within 8–10 weeks of onset of the illness, which is severe and prolonged, although the prognosis is good.

BACTERAEMIA Disseminated infection may occur in association with a local lesion and metastasize to new areas. This may occur with many organisms. As an example, in meningococcal infection, meningitis is not always present. In the early stages, patients are acutely ill, with fever, chills, arthralgia and myalgia, particularly severe in the legs and back. They are very prostrated and hypotensive, and 70 per cent develop a characteristic petechial rash (Fig. P47). Meningococci are cultured from the blood and sometimes from scrapings from the skin lesions, and from the cerebrospinal fluid in cases with meningitis. A rare form of chronic meningococcaemia has been identified that lasts for weeks or months, and this is characterized by fever, rash and arthritis or arthralgia.

Figure P47 Meningococcal septicaemia that presented as subarachnoid haemorrhage and purpuric rash.
**Infective endocarditis**

The comparatively rare acute or malignant endocarditis is due to infection by one of the pyogenic bacteria, such as haemolytic *Streptococcus*, *Pneumococcus*, *Gonococcus* and *Staphylococcus*. Many other organisms may cause endocarditis: *Corynebacterium diphtheriae*, *Brucella* and numerous others have been reported as causative agents. Pseudomonal endocarditis may follow open-heart surgery. The type formerly termed the ‘subacute’ variety occurs in a subject with chronic valvular disease of congenital or rheumatic origin, when the organism is usually *Streptococcus viridans*. Patients present with cardiac symptoms, anaemia, cerebral vascular lesions or, most commonly, pyrexia of unknown origin, a feature that arouses suspicion when progressive anaemia of normocytic and orthochromic type develops. The white cell count is variable, and a polymorphonuclear leucocytosis up to about 9000–12 000 per mm$^3$ is common. Other diagnostic features are an enlarged spleen, clubbing of the fingers in half of the subacute cases (but rarely in the malignant type), petechial haemorrhages under the nails and in the retina and conjunctiva, and Osler’s nodes. Emboli may occur in any organ or tissue. Red blood cells in the urine are invariably found to a greater or lesser degree.

Blood cultures should be performed immediately, as treatment must not be delayed. If over middle age, a patient who is not immediately treated may be cured of the infection but die of cardiac failure due to the damage to the heart.

**LOCALIZED INFECTION**

Many cases of continued fever are due to localized infection (e.g. abscess formation). The elicitation of local signs, in addition to pyrexia and other evidence of generalized systemic illness, will provide the diagnosis, although it may be long delayed.

A rectal or vaginal examination should serve to detect prostatic abscess, periproctal abscess, ischiorectal abscess, pyosalpinx, suppurring ovarian cyst or parametritic abscess, all of which are likely to cause local pain in the perineum, anal region, sacral region, back or lower abdomen.

In its acute form, *empyema of the maxillary antrum* causes pain and tenderness over the affected maxilla, with oedematous swelling of that side of the face, but in chronic cases the symptoms may be much less definite. The diagnosis may be suggested by facial pain, local swelling and perhaps an intermittent purulent discharge from one nostril. Radiographs and antral aspiration will confirm the diagnosis.

*Empyema of a frontal sinus may be acute or chronic, causing pyrexia in either case. The diagnosis may be suggested by the complaint of local headache above the eyes, generally on one or other side of the midline rather than central, and especially if the headache is associated with local tenderness to percussion. Identification becomes easy if the abscess points above the inner canthus of the orbit near the root of the nose, although doubt may persist for a long time. Difficulty in diagnosis applies still more to *empyema of the ethmoidal or sphenoidal sinuses*, in which few objective signs are to be expected. The patient may complain of severe frontal headaches that are often worse in the morning but pass off later in the day. A purulent nasal discharge may also be present, while the pyrexia may be only slight, but generally persistent.*

*Suppurating lymph nodes* will be diagnosed from the character of the tender swellings that precede the skin-reddening and the actual formation of an abscess. The site is likely to be the neck, axilla or groin, and there will usually be an indication of the source of the problem in the form of a septic focus in the skin corresponding to the lymph drainage of the node concerned – impetigo, a septic cut, a wound or a whitlow. One source of trouble that may be overlooked is *pediculosis* of the scalp; this should be suspected if there is irritation of the back of the neck at the roots of the hair in association with enlargement of the occipital as well as the cervical lymph nodes.

*Mammary and submammary abscess* may be of the chronic type and cause pyrexia without much pain. *Empyema thoracis* is generally easy to diagnose. The abnormal physical signs at the base of one lung suggest the presence of fluid, while pus may be found on needle aspiration. The condition may be simulated by subdiaphragmatic abscess, but X-ray examination or ultrasonography will usually help in distinguishing between the two. In some cases, both conditions are present. Difficulty in diagnosis may on occasion be considerable when the empyema is interlobar, or between the pericardium and the pleura, or between the diaphragm and the lower lobe.

*Lung abscess* may be either single or multiple, and may be part of a blood-borne or local infection, or be associated with a bronchial neoplasm (Fig. P48). In the latter situation, much of the fever and acute systemic upset is due to infection rather than to the primary neoplasm. Imaging will help in the differential diagnosis.

*Hepatic abscess*, especially amoebic, may be a subacute or chronic rather than an acute condition,
with fluctuating pyrexia persisting for months. The diagnosis may be suggested by a complaint of pain or tenderness over the lower part of the right chest in front or behind, by dullness at the base of the right lung, or by friction sounds over the liver. A history of amoebic dysentery is not always forthcoming. Pain is in some instances referred to the right shoulder. When pyrexia and rigors are the only objective features, malaria is simulated, but a high polymorphonuclear leucocytosis is against this diagnosis. Elevation of alkaline phosphatase may be the only clue in some cases. The diagnosis of hepatic abscess is clinched by ultrasound-guided needle aspiration, which reveals pus, or fluid resembling ‘anchovy sauce’ in amoebic abscess. On rare occasions, this may be coughed up as the result of ulceration through the diaphragm and pleura into a bronchus.

In empyema of the gallbladder, jaundice is generally absent. The pyrexia may be considerable and prolonged, and rigors are to be expected. The diagnosis depends largely upon the patient’s complaint of pain in the right hypochondrium, associated either with an enlarged gallbladder or with acute pain and tenderness on palpation of the gallbladder region below the tip of the right ninth rib cartilage.

Suppurative cholangitis is the result of the extension of pyogenic infection up the hepatic ducts into the biliary canals within the liver. It generally is associated with obstruction in the bile duct by stone or growth. By the time the infection has extended to become suppurative cholangitis, the patient will have become increasingly ill. The supervention of cholangitis may be indicated by progressive, soft, uniform and tender enlargement of the liver, associated as a rule with jaundice. Recurrent rigors are almost invariable.

Suppurative pylephlebitis arises from infection somewhere in the periphery of the portal area – for example, previous appendicitis. The condition is often fatal. The liver becomes studded with multiple small abscesses around the intrahepatic subdivisions of the portal vein. The liver becomes progressively, smoothly and uniformly enlarged, and generally tender; jaundice is present in fewer than half the cases. The high degree of pyrexia, the rigors, the asthenia and the wasting all indicate that the patient has developed some form of septic extension of the original disease. There is a high degree of leucocytosis.

Subdiaphragmatic abscess is often difficult to diagnose. There may be no abnormal physical signs, but more usually infection of the pleura through the diaphragm leads to an impaired percussion note at the base of one lung, accompanied by pleuritic friction and rales. Diagnosis is aided by the use of ultrasonography or computed tomography, followed by needle aspiration.

Bronchiectasis may be responsible for prolonged periods of pyrexia with afebrile intervals of varying length. The pyrexial bouts are due either to invasion of the pus-containing cavities by fresh organisms, or to recrudescence of infection already present, possibly brought about by impaired bronchial drainage. The abnormal physical signs in the lungs, the abundant foul green sputum and the clubbed fingers indicate the diagnosis. Not infrequently, there is considerable inflammation in the lung tissues around the area of bronchiectasis – the so-called peribronchiectatic pneumonia. This condition should be suspected in any patient in whom consolidation, often with pleurisy, recurs repeatedly in one area of the lung.

An appendix abscess may be easy to diagnose on palpating the tender swelling in the right iliac fossa. In other locations, the abscess may be difficult to diagnose, a rectal examination leads to the detection of the abscess when it descends into the pelvis. Pyrexia ceases as a rule when the pus obtains free drainage, so that it is exceptional for appendix abscess to be the cause of prolonged pyrexia.

Perinephric abscess may cause pyrexia of considerable degree, possibly lasting for several weeks. Pain in the loin is almost always present, eventually with tenderness to palpation in both the loin and the lumbar
region, but it is often absent in the early stages and may not appear until the patient has been febrile for some weeks. There may be no defined swelling, but only a sense of resistance evident when, with the patient recumbent, the examiner places one hand behind each loin with the fingertips external to the erector spinae muscles, and then makes as if to raise the patient from the bed, although without actually lifting. The fingers on the affected side will not feel the hollow of the loin as clearly as will those on the sound side. The signs may be yet more striking if the patient lies prone. If they are well enough to sit up in bed with the back bared, the observer may then look down the patient’s spine from above. It may often be apparent that the loin on the sound side is slightly concave, while that of the perinephric abscess side is either flat or slightly convex. Only in pronounced cases does the loin show a distinct convexity. Perinephric abscess is generally the result of pyogenic infection within the kidney; alternatively, it may be due to pus tracking up behind the colon from appendicitis, or it may be a delayed result of a loin injury, a haematoma due to the injury becoming infected and slowly forming a perinephric abscess weeks or months after the trauma. In many cases, the history is obtainable of a suppurative process a short time previously. Pre-existing neutropenia or corticosteroid therapy will in these cases – as in all infective processes – predispose to abscess formation.

**Diverticular abscess** may be subacute and yet cause prolonged pyrexia. It is generally situated in the left lower part of the abdomen, producing a tender swelling that may simulate carcinoma. It is preceded by chronic bowel symptoms, constipation and colic. Bleeding may occur, sometimes profusely. It is a disease of the second half of life.

**Psoas abscess** results from tuberculous spinal disease; the condition may be apyrexial but, like any other form of tuberculosis, it may cause protracted irregular pyrexia. Pain localized to some part of the back and stiffness of the corresponding part of the spine in a child are suggestive features. Radiographs must be taken. On the other hand, the diagnosis may remain unsuspected until a tender swelling appears above or below one groin as the abscess tracks down from the spine along the course of the psoas muscle, ultimately causing fluctuation from above to below the inguinal ligament.

**Actinomycosis** is diagnosed by the discovery of the organism. It may be in the discharge from a sinus communicating with the focus infected, generally the cheek, jaw, neck, lung, liver, caecum or spine. It may, however, occur anywhere in the skin or viscer, and the disease is likely to be missed if specific bacterial investigation is not undertaken. An actinomycotic ischiorectal abscess, for instance, may be regarded as of merely pyogenic origin. There is diffuse infiltration of deep as well as superficial parts, a liability to discharge through one or more sinuses, and a suggestive purplish-red colour of the skin adherent to the lesion. The course is chronic and often apyrexial, but frequently there are periods of pyrexia.

**INFECTIVE AND INFLAMMATORY CONDITIONS**

*Coliform infection* of the urinary tract may be chronic and apyrexial, but it is liable to exacerbations with prolonged pyrexia, aching or pain in one or both loins, frequency of micturition, and pain during micturition. It may exist, however, especially in children, with so few symptoms of urinary disease that its responsibility for continued pyrexia may be missed.

Chronic or recurring pyelonephritis is more common in women than men, but it may be associated with an enlarged prostate or urethral stricture. The patient is ill, with rigors and high, long-continued pyrexia; the urine is purulent, and yields a positive culture of the causative organism or organisms. In papillary necrosis, the renal calices become clubbed and pyuria is common, even when urine cultures are sterile.

Prolonged fever is not uncommon. It may be due to prolonged taking of compound analgesic tablets or diabetes mellitus.

**Gallstones** may be silent, causing no symptoms, or they may be associated with irregular and sometimes prolonged pyrexia, and with bouts of pyrexia in attacks of biliary colic.

**Phlebitis** in a superficial vein is indicated by tenderness, with or without redness and swelling along the course of the vein; pyrexia of variable degree and duration accompanies the disorder in the earlier stages but usually subsides in a few days. The diagnosis is much more difficult when the inflamed vein is deeply situated. Intra-abdominal phlebitis may be responsible for both continued pyrexia and vague, but possibly severe, abdominal pain in certain cases for which no explanation is forthcoming. *Thrombophlebitis migrans* occurs uncommonly. Venous thrombosis in the pelvis, not necessarily associated with obvious femoral or popliteal thrombosis, may account for pyrexia after childbirth.

**Thyroiditis**, an inflammatory but non-infective condition, may cause the complaint of sore throat, the thyroid itself being painful and tender.

**Parametritis** is diagnosed by pelvic examination. It is likely to be the after-effect of recent labour, and is often associated with continued pyrexia, and pain in the
pelvis and lower part of the back. Abscess formation may occur. Elderly women are apt to develop a purulent form of endometritis, sometimes pyrexial, with pelvic pain, bearing-down pain, pain in the back, and a foul vaginal discharge that is often blood-stained, the condition simulating advanced carcinoma of the body of the uterus.

Vesiculitis, although perhaps of local origin, is generally due to a gonococcal infection of the seminal vesicles. The complaint is mainly of hot burning pain in the rectum, aggravated by defecation; proctitis or carcinoma of the rectum or acute prostatitis is simulated. Diagnosis is established by rectal examination, the finger locating a tender swelling in the vesicles.

Colitis, whether infective or ulcerative, will be suggested by a history of diarrhoea, with the passage of blood and mucus associated with more or less pain along the course of the colon, particularly the descending colon; carcinoma or diverticulitis may be simulated. The diagnosis is confirmed by endoscopy, barium enema and/or bacteriological studies.

Crohn’s disease (regional enteritis) should be suspected when there is a history of chronic intermittent diarrhoea, fever, loss of weight and abdominal pains or distension. Barium studies are necessary. Intermittent small-bowel obstruction is common. Pancreatitis, when subacute or chronic, may be very difficult to diagnose. It is sometimes (but not always) pyrexial, and it may simulate other abdominal lesions such as gallstones. Glycosuria in association with pyrexia and a dull aching pain in the abdomen across the site of the pancreas may be suggestive, but the symptoms are generally too vague to be characteristic. There is often a curious dull-brown pigmentation of the skin. Chronic pancreatitis should be suspected in a patient with recurrent abdominal pain, particularly if the pain or tenderness extends to the left of the midline, if gallstones are present, and if there have been bouts of overconsumption of alcohol. Radiographs may show pancreatic calcifications. Repeated serum amylase estimations taken within 12 hours of an acute episode are elevated in most cases, but as more acinar and ductal cells are destroyed, these become less evident. Following acute pancreatitis, suppurative pancreatitis may occur in the second or third week with a return of fever.

Familial Mediterranean fever (paroxysmal polyserositis or periodic fever) is an inherited disease of unknown aetiology that is characterized by acute episodes of self-limited fever with signs of inflammation of the peritoneum, pleura and joints. The febrile episodes recur irregularly and unpredictably. The disease occurs most commonly in patients of Mediterranean or Middle East origin, particularly in Sephardic (but not Ashkenazic) Jews, Armenians, Turks, Arabs, Greeks and, less commonly, Italians and others. In the acute form, attacks of fever may reach 40 °C (104 °F), but symptoms of peritonitis or ‘pleurisy’ usually subside within 48 hours. Small pleural effusions may occur. Abdominal symptoms occur most often, occurring in over 95 per cent of patients, sometimes mild, but sometimes with severe localized pain that spreads over the whole abdomen, associated with abdominal distension and muscle rigidity and sometimes ileus, so that an acute surgical emergency may be suspected. Acute arthritis is less common, usually affecting one joint, the knee in most cases. Such arthritic episodes usually last for only a few days, but they may occasionally last for weeks or even months. The onset of this disorder is usually in childhood or adolescence, although it may come on at any age, with males being affected more often than females. Amyloidosis is a complication commonly seen in Israel, and less commonly in the USA. The prognosis depends on the development, or absence, of amyloidosis. In about 25 per cent of cases, transient inflammatory skin lesions like erysipelas occur, usually below the knees.

Sarcoidosis, a chronic granulomatous inflammatory condition, may cause prolonged fever, sometimes with relatively little systemic upset. Hilar glandular enlargement on X-ray examination, erythema nodosum, and a weakly positive or negative tuberculin test are suggestive findings. Diagnosis may be achieved by bronchial and transbronchial biopsy performed at bronchoscopy in cases with an abnormal chest X-ray.

**NON-PURULENT HEPATIC CONDITIONS**

Quite apart from fever that occurs in obviously infective lesions of the liver, such as hepatic abscess, acute viral hepatitis, cholangitis and pylephlebitis, pyrexia – generally without the ordinary concomitants of fever – often occurs when the liver tissue is affected by lesions that are not obviously pyogenic, particularly cirrhosis and carcinoma. Laennec’s (alcoholic) cirrhosis, post-necrotic and biliary cirrhosis may all be accompanied by prolonged fever, as may chronic active hepatitis, a disorder induced by the hepatitis B or C virus, by autoimmune aetiology, or precipitated by various drugs including methyldopa, isoniazid and nitrofurantoin. Hypergammaglobulinaemia is a striking feature. Liver biopsy is necessary to confirm the diagnosis. Rapidly growing neoplasms are sometimes accompanied by pyrexia. The appetite may be fairly
good, and the patient may even be carrying on their ordinary work, although their health is failing.

CONNECTIVE TISSUE DISORDERS
In some cases, prolonged fever is a part of the clinical picture of rheumatoid arthritis. Systemic lupus erythematosus may for many months present as pyrexia, often accompanied by skin rashes and symptoms relating to locomotor and other tissues. The high erythrocyte sedimentation rate and joint pains may lead to confusion with rheumatoid arthritis. Polyarteritis nodosa and other vasculitidities, such as giant-cell arthritis or Wegener’s granulomatosis, now referred to as granulomatous polyangiitis, may cause prolonged fever, although the latter is often rapidly fatal in a few weeks, or sometimes less. Polymyositis and dermatomyositis may also be associated with fever, as may (although rarely) systemic sclerosis (scleroderma).

BLOOD DISEASES
Any one of the severe blood diseases may be associated with prolonged pyrexia. Diagnosis depends upon other factors, particularly the blood count, as in leukaemia. In agranulocytosis and aplastic anaemia, the infection is responsible for the fever, but in any severe prolonged anaemia – particularly in childhood – prolonged pyrexia may be seen. In previous times, for example, untreated Addisonian anaemia was a febrile disease. Febrile episodes occur also in haemolytic anemias.

DISEASES OF THE TROPICS AND SUBTROPICS
Trypanosomiasis (sleeping sickness)
Trypanosomiasis is a parasitic infection that occurs particularly in Africa where Glossina palpalis, the tsetse fly, abounds. The bite of this insect spreads the disease – sleeping sickness – by invasion of the central nervous system. It is not always pyrexial, and at times malaria may be simulated. The trypanosome may be identified in blood films, lymph node aspirates, more rarely bone marrow or, in the final sleeping stage, the cerebrospinal fluid. There is no distinctive feature on the temperature chart.

Malaria
The main types of malaria are: benign tertian (Plasmodium vivax or Plasmodium ovale), in which rigors occur on alternate days with a maximum temperature of 39.4–40 °C (103–104 °F) (Fig. P.49) and complete freedom on the intermediate days;

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Figure P.49 Temperature chart of simple tertian malaria, showing attacks occurring every third day. [London School of Tropical Medicine.]
PYREXIA, PROLONGED

quartan (*Plasmodium malariae*), in which there are 2-day intervals so that the paroxysms occur every fourth day (Fig. P50); and malignant tertian (*Plasmodium falciparum*), in which the fever is often more irregular. The intervals between the attacks of fever vary in accordance with the time that successive generations of the various strains of parasites take to mature. A patient may be infected by one set of bites by a mosquito with a tertian or quartan ague, and subsequently become infected by other mosquitoes with either tertian or quartan parasites. Thus, there would be a mingling together of the effects of different generations of *Plasmodium*, and the patient would have a daily (quotidian) paroxysm (Fig. P51). Similarly, infection by two batches of parasites might result in a complicated clinical picture in which the attacks of pyrexia might be irregular or almost continuous. As a rule, a paroxysm, with its various cold, hot and sweating stages, lasts for about 8 hours. The diagnosis of malaria will be confirmed by the discovery of parasites on analysis of thick films of the blood.

One remarkable feature is that malaria may remain latent for many years, particularly with *Plasmodium malariae*. However, considering the large number of serving men infected during the Second World War, relapses of malaria 1 year after returning home were rare. Reappearance is brought about by a general deterioration in health, or through some intercurrent illness.

Kala-azar (visceral leishmaniasis)

There is no characteristic chart, and the pyrexia is often extreme and of a swinging, but continued, type. Great enlargement of the spleen with continued pyrexia would suggest the diagnosis, which is confirmed by discovering Leishman–Donovan bodies from material obtained by splenic or sternal puncture. This condition was not uncommon on the Mediterranean coast, and even today may be acquired on a short continental holiday, albeit rarely.

Plague and cholera

Plague is also epidemic, the two best known types being the bubonic and the pneumonic. The diagnosis depends on discovering plague bacilli (*Pasteurella pestis*) in fluid obtained by aspirating a bubo, or from the sputum by special bacteriological methods. The pneumonic form is the more acute, and it may simulate lobar pneumonia; the bubonic form is of longer duration, with a lower grade of pyrexia.

Both cholera and plague, when they occur in the UK, tend to be in persons coming from India or the East, particularly members of ships’ crews.

Relapsing fever

At one time, this condition was prevalent in Great Britain and it acquired the name “famine fever” from the circumstances of its occurrence. It is due to infection with *Borrelia recurrentis* being introduced by an infected louse when crushed against an abrasion or wound. Many strains are also conveyed by ticks, the disease being transmitted by their bites. The course of the disease is characteristic. There are outbreaks of high pyrexia (Fig. P52) associated with extreme prostration and severe illness lasting 5–6 days, alternating with complete intermission of about the same duration. There may be an indefinite number of relapses before death or recovery. The spirochaete may be identified in a blood film shortly before the febrile paroxysm but not in the intervals.

Schistosomiasis

This condition is widely spread through Egypt, Zimbabwe and certain other parts of Africa. Involvement of the urinary tract with *Schistosoma haematobium* is extremely common in these countries. *Schistosoma mansoni* affects the colon more and is associated with splenomegaly. Pyrexia over long periods is not uncommon with either type of infection.

MENINGEAL AND CEREBRAL HAEMORRHAGE

Almost any pathology that interferes radically with the heat-regulating centres in the brain may cause hyperthermia, often without the other symptoms usually associated with fever. As a rule, death or recovery renders such types of pyrexia of short duration, for instance in cases of pontine haemorrhage. Sometimes, however, the patient may survive long enough to come into the category of cases of prolonged pyrexia.

SKIN CONDITIONS

These are discussed in detail elsewhere (see BULLAE AND VESICLES (BLISTERS), p. 76). There is less tendency to pyrexia in dermatitis herpetiformis, herpes gestations and erythema iris than in acute and subacute pemphigus. Pemphigus is always a serious and often a fatal malady, the skin blebs being but a local manifestation of some severe but ill-defined systemic reaction. Eosinophilia is usual. Any diffuse inflammation of the skin, from acute psoriasis to a gold dermatitis, may be accompanied by fever.

MALIGNANCY

Lymphomas of different types, including Hodgkin’s disease, may be associated with prolonged fever, the
**Figure P.50** Temperature chart of quartan malarial fever, the attacks recurring every fourth day. [London School of Tropical Medicine.]
Pel–Ebstein type depicted in Figure P.53 being seen typically in this condition, although not confined to it. Lymphosarcoma may be associated with prolonged pyrexia, as may carcinoma, particularly if associated with sepsis, as is seen not infrequently in the bronchus and large intestine. Infection is not necessary for pyrexia to be present; high fevers are occasionally seen in primary and secondary carcinoma where no sepsis exists, and a low-grade pyrexia is common. Particularly likely to be accompanied by prolonged pyrexia are hypernephroma, hepatoma, secondary carcinoma in the liver and widespread metastatic disease.

ALLERGIC (ANTIGEN–ANTIBODY REACTIVE) CONDITIONS

Being inflammatory disorders, these are often pyrexial. This is seen, for example, in Henoch–Schönlein syndrome (anaphylactoid purpura) and various skin allergies. The post-cardiac injury syndrome may occur after any injury to the heart, whether it is an operation, stab wound or myocardial infarction (Dressler’s syndrome). Fever, pericarditis, pleurisy and pneumonitis may occur 2 weeks or more after the injury, and recurrences are common.

FACTITIOUS PYREXIA PRODUCED BY MALINGERERS

From time to time, there arise cases in which illness is simulated by a patient who deliberately deceives by one or other of various devices, such as dipping the thermometer bulb in a cup of tea, holding it against a hot-water bottle or rubbing it violently against the blankets. It is said that some have the art of squeezing the bulb between the fingers or between the teeth with just sufficient force to raise the temperature, but without breaking the glass. On suspicion of such a possibility, the temperature is taken – preferably per rectum – under careful scrutiny. If a factitious pyrexia is suspected, the patient may be left unattended on the first occasion; if the temperature is raised, it should then be taken again immediately under close observation. A marked difference between the two readings strongly suggests that the first reading has been faked. Cases are also recorded where patients have deliberately infected themselves to produce a fever.

DRUG REACTIONS

In any persistent unexplained fever where drug treatment is being given, the possibility of the therapy...
**Figure P.52** Four-hourly temperature chart in a case of severe relapsing fever.
Figure P.53 Morning and evening temperature charts in a case of Hodgkin's disease, illustrating the Pel–Ebstein type of recurrent or periodic pyrexia that may occur in this disease.
being the cause should be considered. Fever occurring in the treatment of pulmonary tuberculosis, for example, may be due to isoniazid, aminosalicylate or streptomycin, or to all three drugs. Antibiotics, sulphonamides, arsenicals, iodides, barbiturates and many others may be responsible. When in doubt, the motto should be, if possible, ‘Stop treatment’.

PYREXIA, WITHOUT OBVIOUS CAUSE
Mark Kinirons

FEVER IN CHILDREN
Temperature-regulating reflexes may be underdeveloped or poorly controlled in children and infants. Body temperature is apt to be disturbed in children by causes that would not produce pyrexia in an adult; hence a transient, irregular or recurrent rises of temperature in a child may often be of little significance. Immediately after birth, temperatures of around 38 °C (100.4 °F) are not uncommon and are physiological, but fever may persist with birth injuries associated with intracranial bleeding.

Nevertheless, if such rises of temperature persist, it is unwise to regard them as purely trivial. There are three conditions in particular that need to be specially borne in mind: (i) urinary infections; (ii) upper respiratory tract infections; and (iii) exanthemas and the infectious diseases of childhood before their specific rashes or other clinical features appear.

Urinary infection
Usually due to Escherichia coli, this is only too easily missed in a child, as there may be no symptoms attracting attention to the urinary system. The presence in the urine of leucocytes makes the diagnosis likely, although it needs culture of a clean specimen of urine to establish it.

Upper respiratory tract infections
These include tonsillitis, acute pharyngitis and rhinitis, and otitis media. Although symptoms usually point to the cause of the fever, in a small child this is not always so; the child has fever, is distressed and apparently in pain, but they cannot indicate to the parents where the trouble lies.

Of the many other causes of fever in childhood, starvation, dehydration and acidosis should be mentioned. Tuberculosis in the lung, lymph nodes or elsewhere is much less common now in developed countries than previously, but it is still common in many parts of the world. The only sign of tuberculous disease initially in a child may be unexplained fever, but the same is true of a large number of infective processes. Of course, bacterial meningitis should always be borne in mind in a listless child with fever. Evidence of meningism, photophobia and skin lesions should be sought.

PYREXIA IN ADULTS
The types of obscure pyrexia in adults fall into two main groups: (i) transient or short pyrexia, and (ii) continued pyrexia.

Transient or short pyrexia
This is usually due to one of a number of mild viral or bacterial infections, the common cold, sinusitis, nasopharyngitis, mild pyelonephritis and a host of others. It is as well to remember that drugs may themselves cause fever, and that the curative drug may sometimes maintain the fever it was used to cure. It is also as well to remember that products of tissue destruction may cause fever, as after myocardial infarction, pulmonary infarction, gangrene and tissue necrosis, and accumulation of blood in body cavities, gut, joints or elsewhere.

Continued pyrexia
This may occur in any of the large number of conditions listed under PYREXIA, PROLONGED (p. 558). In Great Britain today, one thinks primarily of the more common conditions: coliform and other infections of the urinary tract, infectious mononucleosis, carcinoma, Hodgkin’s disease, non-Hodgkin’s lymphoma, leukaemia, tuberculous disease, chronic sepsis (as in sinusitis), abortus infections and so on. Fever may also be associated with AIDS with its intercurrent infections. With air travel readily available, malaria, kala-azar and any other tropical disease may be imported overnight and, with a large immigrant population from overseas, conditions such as amoebiasis, schistosomiasis and leprosy may be encountered in the ordinary outpatient clinic.

FEVER IN THE AGED AND DEBILITATED
Febrile disorders are often missed in the elderly, the debilitated and immunosuppressed individuals (such as those receiving steroids), as infections may be accompanied by much less fever than usual. In some cases, absence of fever in the presence of acute infection carries a bad prognosis.

PYURIA
Ben Challacombe

Pyuria means no more than the presence of pus/leucocytes in the urine. It will be present when there are infective processes affecting the urinary tract,
in some chronic non-infective conditions such as bladder carcinoma-in-situ and interstitial cystitis, and occasionally following the rupture of an abscess outside the urinary tract into the system. The quantity of pus may vary; when present in large quantities, it forms a thick grey or yellow sediment.

On microscopy, pus cells (often neutrophils) are seen as rounded multinucleated bodies about twice the size of a red cell. As pus cells are, in fact, protein, dipstick testing is almost invariably positive for this substance. This test will also be positive if there are abnormally large numbers of epithelial cells in the sample so that microscopy of a urine sample is the only reliable simple test for the presence of pus.

The site of the pus-producing lesion cannot be determined simply by examination of the urine. The general and specific history of the individual case is essential, although, in general, vesical lesions are often consequent upon renal lesions, particularly when these are infective. For example, when a pyelonephritis arises as a result of haematogenous spread, the initial symptoms may be those of cystitis. In time, the pyrexia, rigors and sweating attacks of pyelonephritis become manifest, together with severe loin pain on the affected side. A pyonephrosis need not necessarily present with pyuria if outflow from the kidney is blocked for one reason or another, such as a stone.

**SPECIAL INVESTIGATIONS**

**Imaging**

If the patient is investigated during an acute illness associated with pyuria, imaging investigations of the urinary tract may provide specific answers with regard to the site of the pus. The kidney that is the origin of acute pyelitis/pyelonephritis will appear larger on ultrasound examination, and the fluid in the collecting system will appear less transonic than normal urine, implying turbidity. Complete cessation of renal function is occasionally observed in acute interstitial bacterial nephritis.

**Cystoscopy**

Instrumentation of the lower urinary tract is not recommended during an acute illness as the risk of sepsicaemia is considerable. However, following appropriate antibiotic therapy and the resolution of gross infection, cystoscopy under further intravenous antibiotic cover may provide useful information.

It is most likely that the bladder will be the source of pyuria if it shows the obvious features of a cystitis, but if the mucosa is normal on cystoscopy, inspection of the two ureteric orifices may yield valuable information:

- A ureterocele is seen as a bulge above and lateral to the ureteric orifice, disappearing as the pressure inside the bladder increases with filling.
- A refluxing ureter is wider than it should be and is often placed far laterally. The ureteric efflux may be thick and turbid, and exuding like toothpaste from the orifice if the urine production from that side is diminished.
- The ureteric orifice may be oedematous if pyelitis has extended along the whole length of the ureter as secondary ureteritis.
- The ureteric orifice in chronic tuberculosis is characteristically ‘golf-hole’ in appearance.
- Tiny nodules resembling sand granules are seen around the orifice in schistosomiasis.
- The areas above and lateral to the orifices are those characteristically affected by early carcinoma-in-situ, looking very much like inflammatory patches.
- Interstitial cystitis (Hunner’s ulcer), presents as a red vertical line on the posterior wall of the bladder. This line extends into a split as the bladder is distended and rivulets of blood are seen on the back wall of the bladder, falling down as a curtain into the area behind the trigone.

A classified list of the causes of pyuria is provided in Box P.18.

**Box P.18 Causes of pyuria**

<table>
<thead>
<tr>
<th>Pyuria from diseases of the urinary organs</th>
<th>Urethral</th>
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</thead>
<tbody>
<tr>
<td>Renal</td>
<td>Diverticula</td>
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<tr>
<td>– Pyelonephritis</td>
<td>Bilharzia haematobia</td>
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<tr>
<td>– Renal abscess</td>
<td>Trichomoniasis</td>
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<tr>
<td>– Renal papillary necrosis</td>
<td>Carcinoma-in-situ</td>
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<tr>
<td>– Pyonephrosis</td>
<td>Interstitial cystitis</td>
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<td>– Tuberculosis</td>
<td>–</td>
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<td>– Calculus</td>
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<tr>
<td>– Medullary sponge kidney</td>
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<tr>
<td>– Calculus</td>
<td>Urethral</td>
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<td>– Megaueterter</td>
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<td>– Ureteric foreign body</td>
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<td>– Vesicooureteric reflux</td>
<td>–</td>
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<tr>
<td>Vesical</td>
<td>–</td>
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<tr>
<td>– Cystitis</td>
<td>Prostatic</td>
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<tr>
<td>– Tuberculosis</td>
<td>–</td>
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<tr>
<td>– Calculus or foreign body</td>
<td>Prostatic abscess</td>
</tr>
<tr>
<td>– Ulcer – simple, epitheliomatous</td>
<td>Calculus</td>
</tr>
</tbody>
</table>

(Continued)
Urinary tract pyuria

The kidneys

Recurrent attacks of pyelitis/pyelonephritis are most common in childhood, during pregnancy and after childbirth. In childhood, pyelitis secondary to reflux often remains undetected and therefore untreated; chronic pyelonephritis in later life is a serious complication.

The kidneys may be infected:
- **Through the bloodstream**
- **By direct spread from the bladder** – which can occur if there is vesicouretic reflux
- **By infections ascending through peri-ureteric lymphatics** – from an infected bladder or from a para-vascular structure
- **By direct spread from the bladder** – along submucosal planes

**Outflow tract obstruction** will predispose to ascending infection, particularly if the bladder detrusor pressure is raised in order to expel urine. This explains the occasional association of pyelitis with benign prostatic hyperplasia, and almost certainly accounts for the increase of incidence of pyelitis in pregnancy. It is unusual for both kidneys to be affected simultaneously, but it is certainly true to say that, once a kidney is damaged, it is more likely to be a seat of infection on subsequent occasions.

**Congenital abnormalities** in themselves will not predispose towards sepsis; however, if there is interference with free drainage in a horseshoe kidney (Fig. P54), an ectopic kidney or a kidney with a degree of pelvi-ureteric junction obstruction, it may be the seat of infection more frequently than a normal system. A duplex kidney may become infected in one or other of its moieties.

**Vesicoureteric reflux** arises when the intramural course of the ureter is short and the normally acute ‘ureterovesical angle’ is lost. The normal ‘hydraulic valve’ that prevents reflux is not therefore present, and there is no obstruction to the retrograde passage of urine. At the same time, acute cystitis may in itself alter the efficiency of this ureterovesical angle so that reflux can occur secondary to the primary cystitis.

Cystoscopy of a patient with reflux often shows a small saccule or shallow diverticulum above and lateral to the ureteric orifice. It seems more likely that this is part of the original maldevelopment than a traction diverticulum.

**Megaureter in children** is a condition in which the whole of the ureter may be dilated, although the condition more commonly affects the distal ureter only. A megaureter may be obstructed or non-obstructed, and may or may not reflux. Paradoxically, an obstructed megaureter may also reflux, with relative obstruction to the passage of urine from upper tract to bladder but a facilitated passage from lower tract to upper.

Megaureter is physiological during pregnancy, either as a result of direct obstruction from the gravid uterus or secondary to the progestogen effect of a maintained pregnancy. Whatever the aetiology of megaureter, the resulting stasis predisposes to sepsis, stone formation, squamous metaplasia, and so on. Reflux may occur in severe cases of outflow tract obstruction where the intravesical pressure rises to a level that overcomes the resistance of the most normal ureteric orifice.

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**Box P.18 (Continued) Causes of pyuria**

- **Vesicular**
  - Seminal vesiculitis, acute or chronic vesicular abscess
- **Pyuria from diseases outside the urinary organs**
  - Leucorhoea
  - Balanitis with phimosis
  - From the extension of inflammatory processes to the bladder, or rupture into the bladder or urethra of an abscess, such as:
    - Prostatic abscess
    - Appendicular abscess
  - Iliac or pelvic abscess
  - Abscess due to colonic diverticulitis
  - Psoas abscess
  - Pyosalpinx
  - Carcinoma of the uterus, rectum, caecum, sigmoid or pelvic colon
  - Ulceration of the small intestine, tuberculous or dysenteric
Reflux is demonstrated by a micturating cystogram, by a MAG3 renogram, and also by ultrasound assessment. Reflux is sometimes severe enough to cause gross distension of the upper urinary tract, especially when there is congenital outflow obstruction, as in urethral valves.

The congenital megaureter–megacystis syndrome arises because of incorrect development of the urinary tract, and the ureteric orifices are widely patent (Fig. P.55). A similar picture may be seen in severe chronic retention when dilatation of the bladder and ureters has occurred beyond the point of recovery when the obstruction is relieved.

**Pyelitis (Pyelonephritis)**

Haematogenous pyelitis arising as an entity separate from other urinary tract pathology is not unusual following acute febrile illness or secondary to supputation elsewhere in the body. The organisms responsible are usually *Escherichia coli*, *Proteus* and *Klebsiella*; *Pseudomonas* is common as a hospital-based infection, as is the increasingly common multiply resistant *Staphylococcus aureus*.

The symptoms of acute pyelonephritis are severe. There are loin pain, symptoms of an associated acute cystitis, hyperpyrexia, tachycardia, rigors and sweating. The urine is turbid and opalescent and may be bright red from haematuria and positive for protein, while microscopy reveals large numbers of free bacteria with pus cells and red cells. The patient may be oliguric because of fluid loss.

While examination reveals a high white cell count and erythrocyte sedimentation rate, urography may show reduced function on the affected side; calculi will almost always show up as radio-opaque bodies overlying the urinary tract.

Chronic pyelonephritis is dangerous because there may be few symptoms early in the course of the disease. A slight stinging during micturition and vague pain in the renal areas may be ignored, and general malaise, lassitude secondary to anaemia, a low-grade persisting pyrexia and hypertension are often presenting features. The urine classically has a specific gravity of 1.010, the blood urea and plasma creatinine are raised, and creatinine clearance is diminished. Ultrasound will show a small shrunken scared kidney. Excretion urography may be impractical because of poor renal function, but delayed films will show irregularity and blunting of the calyces, cortical atrophy and a reduction in renal size. The physical signs of hypertension may be manifest.

**Renal abscess**

Renal abscesses follow acute haematogenous infections and are initially situated in the peripheral area of the cortex. There are general symptoms and signs of a systemic abscess, with acute tenderness in the loin, developing into a mass with an overlying hyperaemic skin. It is rare for renal abscesses to discharge spontaneously through the skin, as they are usually seen and treated well before this happens, but this can occur with prolonged renal obstruction. Occasionally, they may discharge into nearby viscera, the ascending or descending colon and the second and third parts of the duodenum, depending on the side affected. It is uncommon for a left-sided abscess to involve the tail of the pancreas. An abscess may occasionally follow a renal infarct or be related to HIV infection.

**Renal papillary necrosis**

Renal papillary necrosis is common in diabetics. It is now rarely seen following phenacetin abuse as the substance was withdrawn following the discovery of the association between its abuse and renal failure. The renal papillae undergo avascular necrosis and separate from the kidney. They may be passed as sloughs,
in which case the patient may present with ureteric colic, or be retained within the pelvicalyceal system, where they calcify. An acute bacterial infection often supervenes, and presentation with an acute pyelitis or cystitis is not unusual.

Urography shows several changes depending on the severity of the process. In the early stage, a line of contrast can be seen crossing the base of each papilla; as the papilla sloughs, it may be seen as a filling defect within a dilated calyx. Later still, a triangular zone of calcification lying within the pelvicalyceal system is readily apparent. Bilateral renal papillary necrosis can lead to progressive renal failure and death from uraemia.

**Pyonephrosis (Infected hydronephrosis)**

In pyonephrosis, the urine within an obstructed pelvicalyceal system becomes infected, usually secondary to congenital pelvi-ureteric junction or impaction of a stone at the junction. Obstruction of a ureteric orifice by a stone or tumour, or ureteric involvement from a primary bladder or primary uterine carcinoma, is a less common cause of pyonephrosis. The symptoms are not as severe as those of an acute pyelonephritis and are more gradual in their onset. Examination almost invariably shows tenderness in the affected loin, together with a palpable renal mass. Radiological examination often reveals the presence of a stone, while renal function is rarely preserved. The quickest way of proving the diagnosis is by establishing a nephrostomy under local anaesthesia and ultrasound control or inserting a ureteric stent if nephrostomy is not available; this will not only allow aspiration of pus for diagnostic and therapeutic purposes but also provide drainage of the system. Nephrectomy is usually required together with appropriate management of the precipitating cause.

**Renal tuberculosis**

This disease – once common – is now relatively rare in the Western world, and there must always be the danger that the diagnosis will be missed unless it is remembered that all cases of persistent sterile and acid pyuria must be considered as tuberculosis until disproved by the examination of no fewer than three early morning urine samples. Even this number may be insufficient to exclude the diagnosis with certainty, and now a serum PCR blood test is often used. Culture, on specially reinforced media, must be continued for 6 weeks.

The miliary form of tuberculosis that was once seen in childhood is now extremely rare; it is not associated with urinary symptoms. Renal tuberculosis, however, is still very much a reality, the kidney at first being attacked by a tuberculous infection on a microscopic basis. The resultant small tuberculous nodules eventually coalesce to form an area of caseation, which then bursts into the renal pelvis by direct ulceration into a calyx. The transitional cell lining of the pelvis and ureter are subsequently infected with tubercle bacilli, becoming thickened by submucosal infiltration and by oedema.

The symptoms prior to discharge into the urinary tract may be very slight. Aching in the loin may be the only symptom, and albuminuria the only finding once the septic focus has discharged. The symptoms mimic a low-grade pyelitis and cystitis; there is increased aching in the renal area, while frequency of micturition, discomfort while passing urine and polyuria also occur.

The urine is pale, acid, of low specific gravity and turbid; tubercle bacilli may sometimes be found after appropriate staining of a centrifuged sample.

Cystoscopy may show areas of oedema within the bladder, and sometimes small tubercles are visible. The ureteric orifice is usually oedematous and pouting into the bladder. The ‘golf-ball’ change is seen in longstanding disease, when fibrosis has caused contraction of the orifice. Digital examination per rectum or per vaginam may reveal thickening of the bladder wall, and pencil-like thickened ureters are occasionally felt as they hook their way into the bladder base.

Urography/CT urogram may show reduced function on one side, together with cavities, ureteric dilatation and a thick-walled bladder. The caseating areas occasionally undergo calcification, these areas being poorly defined, in contrast to the clear-cut margins of a renal calculus. Calcific caseous debris is sometimes seen passing into the renal pelvis may cause intermittent obstruction, the caseating areas occasionally undergoing caseation, these areas being poorly defined, in contrast to the clear-cut margins of a renal calculus. Calcific caseous debris is sometimes seen passing into the renal pelvis by direct ulceration into a calyx. The transitional cell lining of the pelvis and ureter are subsequently infected with tubercle bacilli, becoming thickened by submucosal infiltration and by oedema.

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renal calculi are radio-opaque (Fig. P.56); all stones are shown on ultrasound of the kidneys or on CT scanning.

**Medullary sponge kidney**
This is a congenital abnormality that probably represents a minor form of polycystic kidneys. The condition is not always bilateral, and it is often associated with the unusual physical sign of body hemi-hypertrophy. Unless the family history is known, the condition usually presents as renal angle pain, pyuria, supervening upper tract sepsis and ureteric colic. The stones may erode into the pelvicalycine system, and from there make their way along the ureter. The radiological changes are pathognomonic of the condition, showing the typical ‘bouquet of flowers’.

**Ureteric disease**

**Ureteric calculus**
Most ureteric calculi will pass of their own accord once they have passed the pelvi-ureteric junction. This is the rule in 95 per cent of cases if the stone is less than 5 mm in diameter. The smaller and smoother the stone, the more likely it is to pass. It is probable that acute ureteric colic is the most severe pain known to man (or woman), the severe pain starting on the affected side radiating upwards to the renal area and downwards towards the bladder and into the testicles or labia. It is surprising how many ureteric stones are, in fact, asymptomatic and present as an incidental finding. Complete blockage of a ureter by a calculus is unusual but, if it happens, it can result in a non-functioning and atrophic kidney. The lower down the ureter the stone impacts, the more lower urinary tract symptoms will be manifest; a stone impacting very near to the bladder can exactly mimic an acute cystitis, with supervening pyelitis, except that there is no pyrexia unless the obstructed and retained urine becomes infected. Calci ni are well demonstrated by urinary tract ultrasound; most stones show on plain X-rays; ultrasound will also give an accurate picture of the degree of upper tract dilatation.

**Ureteric foreign bodies**
It might seem inappropriate to discuss foreign bodies within the ureter, but with the increasing frequency of endoscopic stone surgery, iatrogenic foreign bodies are now introduced with increasing regularity. One of the disadvantages of overvigorous laser or ultrasound stone destruction is fragmentation of the probe or fibre tips; these fragments may embed in the mucosa of the ureter and predispose to pyuria.

It should be noted that non-absorbable suture materials should never be used to close surgical incisions in the ureter as they form an excellent nidus for calculus formation.

**Vesical diseases**
Pyuria will be present in any lesion of the bladder that is associated with inflammation. This applies to acute and chronic bacterial infections, parasitic infections, the presence of a stone, primary and secondary malignant disease, squamous metaplasia, leucopla kia and interstitial cystitis.

**Cystitis**
Cystitis may be acute or chronic and, while both forms are usually associated with infection by
a microorganism, a true infective cause is not essential as any process that produces congestion of the bladder will give rise to cystitis.

In acute cystitis, the mucosa of the bladder is oedematous and congested, leading to epithelial desquamation, pyuria, haemorrhage, the development of small abscesses within the mucosa and occasionally areas of ulceration. The changes are sometimes sufficiently severe to cause sloughing of the whole of the mucosa of the bladder, with profuse haemorrhage.

The symptoms of acute cystitis are well known: there is frequency, urgency, dysuria, perineal pain and pain in the suprapubic area, haematuria and pyuria. The diagnosis of an acute bacterial cystitis depends on the positive culture of an infective organism. While the precipitating cause may be evident in some cases, such as lower tract instrumentation or overvigorous intercourse, many cases cannot be attributed to a specific event. The condition is most frequently seen in young women after they become sexually active, but it is also common in perimenopausal women with poor bladder emptying or urethral stenosis due to poor oestrogenisation.

Chronic cystitis may follow an improperly or inadequately treated acute cystitis. While the symptoms are less severe, increased frequency of micturition, pyuria and a persisting alkaline urine are noted. It is often seen with some form of urinary obstruction or retention, and it is sometimes found in cases of urinary incontinence, of whatever cause. An important second diagnosis is to consider tuberculous cystitis.

At cystoscopy, the bladder wall is red, smooth and oedematous; in pyelonephritis, the cystoscopic appearance of the bladder is normal apart from possible modifications in the appearances of the ureteric orifices, the efflux of which is often cloudy as it contains more particulate matter.

Bladder tuberculosis
This is part of a tuberculous infective process affecting the whole of the urinary system, together with the reproductive system of both sexes. Frequency is the predominant presenting symptom, by day and by night, associated with a minor discomfort usually felt at the end of the urethra. A few drops of terminal haematuria are often observed. Pyuria is constant. Vesical calculus and vesical carcinoma, particularly carcinoma-in-situ, present in a similar fashion. Bladder calculi are usually found in older patients with symptoms of outflow tract obstruction or a history of previous lower urinary tract instrumentation and catheterization. Calculi in the bladder often give pain only on movement; haematuria in calculus disease is observed throughout the stream and is often a more regular occurrence than with carcinoma.

The symptoms of bacterial cystitis may supervene in both calculus and carcinoma, accompanied by frequency, urgency and painful micturition by day and night. Vesical carcinoma may be felt per rectum, especially when the patient is thin and the tumour is extensive. The early stages of tuberculous cystitis are characterized cystoscopically by the appearance of greyish tubercles in the submucosal coat of the bladder, particularly around the ureteric orifice. At this stage, frequency of micturition may be the only symptom, but, as the disease progresses, the tubercles enlarge, coalesce and eventually ulcerate, by which time pus and blood will both be present in the urine; tubercle bacilli should show up on special staining. The bladder becomes extremely small, with micturition occurring every 15 minutes or so throughout the 24 hours.

As a general diagnostic rule, any patient with increased frequency of micturition and a sterile acid urine should be considered as suffering from tuberculosis. Several early-morning urine cultures may be required before the clinical suspicion of tuberculosis can be dismissed. Culture, on special media, must be continued for 6 weeks. Tuberculosis within the bladder is often associated with tuberculosis in the Fallopian tubes, vas deferens, epididymis, prostate and seminal vesicles.

Tuberculosis in the bladder usually arises secondarily from an infection in the upper tract. A caseous lesion, however small, in the kidney ruptures and discharges its contents into the renal pelvis, so that the urothelium of the pelvis, the ureter and the bladder become affected in turn. The primary site may similarly be within the epididymis, infection ascending through the vas deferens to affect the bladder, seminal vesicles and prostate. Prostatic tuberculosis is rare but may involve the bladder by direct ulceration. As with simple infective pyelitis, renal tuberculosis may present with lower urinary tract symptoms; the amount of blood present in the urine is usually less than if the bladder is chiefly involved, and blood will, of course, be noted and found throughout the urinary stream. In renal tuberculosis, there may be tenderness in the loin, while the kidney may be more readily palpable than usual, and the distal end of the lower ureter can sometimes be felt on rectal or vaginal examination.

Ultrasound, CT and intravenous urography may provide useful information with regard to the state of the upper tract. Cystoscopy in renal tuberculosis may show pathological changes surrounding one
The affected ureter is primarily oedematous, and the wall is thickened and patulous. Later, the orifice becomes rigid and patent, representing the 'golf-hole', and is drawn up towards the affected kidney by fibrotic shortening of the ureter. The final stage of vesical tuberculosis is a small thick-walled bladder (the 'golf-ball' bladder).

**Bladder calculus**

Calcus in the bladder often presents as simple bacterial cystitis, and in these circumstances there is little that will distinguish cystitis from many other forms except, perhaps, that the urine may be loaded with crystals. An increase in the amount of blood after exercise is noted frequently, while pyuria is usually constant. Haematuria may occur after exercise, as it does in joggers.

The constant symptoms of vesical calculus are frequency and discomfort during the daytime, especially when erect and moving, penile pain after micturition and haematuria. Except for those patients who have had indwelling catheters or complex lower urinary tract surgery, vesical calculus is almost always secondary to sepsis and stasis from outflow obstruction. The management must aim not only to remove the calculus but also to eliminate the source of obstruction. A suspected calculus will almost always show on plain pelvic X-ray or ultrasound. It may not always be possible to determine whether a stone lies within a diverticulum, except by cystoscopy.

Stones that form in the upper tract and pass into the bladder without producing the symptoms of ureteric colic are almost always small enough to pass per urethram, unless the bladder outlet is obstructed. Radiolucent stones within the bladder (e.g. uric acid stones) are unusual as they accumulate calcium once within the viscus and take on a laminated appearance. If a pure uric acid stone is present within the bladder, it will not show on plain X-ray, but it will probably be detected as a filling defect at urography.

**Ulceration of the bladder**

This occurs secondary to chronic cystitis, following traumatic cystoscopy, secondary to a long-standing stone and as a consequence of radiotherapy for pelvic malignancy. Hunner's ulcer (interstitial cystitis) is a disease that is peculiar to women, causing severe frequency, pain on micturition, urgency with incontinence, and occasional haematuria. The diagnosis is established cystoscopically when a vertical ulcer is usually observed on the posterior bladder wall. The ulcer splits as the bladder is distented, resulting in a curtain of blood rivulets falling down the posterior wall. Biopsy of the area, usually taken to exclude malignancy, shows chronic inflammatory changes with a heavy mast cell infiltrate. Calcium encrustation occurs, so that pyuria and calcific debris are often seen. The bladder capacity in this condition is relatively small, and therapy often consists of forcible distension. Tuberculous ulceration, malignant ulceration and ulceration secondary to radiotherapy have similar presenting symptoms of frequency, haematuria, urgency and additional pain at the termination of micturition. The cystoscopic appearances of these different ulcers are not always easy to distinguish, and multiple biopsies are often necessary in order to establish the diagnosis with certainty.

**Malignant ulceration of the bladder**

Malignant ulceration of the bladder and papillary transitional cell carcinoma of the bladder are common conditions, giving rise to irregular macroscopic haematuria, which is often profuse and almost always painless. Well-differentiated tumours tend to protrude into the bladder lumen, supported by a pedicle of rather narrow size, which accounts for the fact that the surface is often necrotic and ulcerated, giving rise to pyuria in conjunction with haematuria. The pathognomonic symptom is painless haematuria, but increased frequency can occur if the tumour(s) is sufficiently large to disturb bladder capacity. Pain is unusual unless secondary infection has occurred. Tumours are often multiple because of the 'field change' that occurs within the whole of the transitional cell lining of the urinary tract – the urothelium.

When tumours are less well differentiated and situated near the ureteric orifices, there may be ureteric obstruction and loin pain secondary to distension of the affected upper tract. Diagnosis can almost always be established preoperatively by a combination of urography and ultrasound examination of the urinary tract. Endoscopic examination of the bladder is conclusive and allows visualization and biopsy.

When the tumours are large, clumps of the frond-like lesions may separate and be present in the urine. Cytological examination of voided samples is relatively unsatisfactory if the tumour is well differentiated, as the cells are barely different from normal bladder epithelial cells. As soon as relative de-differentiation occurs, cytological examination of the urine is a useful diagnostic and monitoring tool.

It is probably true to say that the more poorly differentiated a transitional cell carcinoma, the more solid-looking it becomes. The solid carcinomas are nodular, sessile, often solitary and involve the trigone rather than affecting the lateral walls. Ureteric obstruction is a frequent complication, and early
invasion of the muscle wall occurs. The presence of a mass on bimanual palpation reveals that the tumour is probably beyond the scope of endoscopic resection, while fixity to the pelvis implies inoperability.

**Diverticulum of the bladder**
A bladder diverticulum may give rise to intermittent or persistent and excessive pyuria together with increased frequency, pain and difficulty with micturition. The last symptom relates to the outflow tract obstruction. A common symptom is that bladder emptying is often followed quickly by the need to empty the bladder again; as the bladder ‘empties’, it expels as much urine into the diverticulum as it does through the urethra, so that when the sphincter apparatus has closed, urine flows back into the bladder cavity from the distended diverticulum. The diagnosis is established by endoscopic examination but may be seen on ultrasound and CT.

**Schistosomiasis**
This causes pus in the urine when the small submucosal nodules – the ‘sandy’ patches – ulcerate into the bladder. Ova are often observed in the urine, together with pus and blood, but microscopic examination of a biopsy is pathognomonic. Complement fixation testing is specific. In advanced cases, calcification, appearing as a ring in the bladder wall, may show on plain X-ray, while urography shows upper tract dilatation, due to the presence of ureterovesical stricture. Complication of the disease process by carcinoma is all too common, and it is to some extent related to the duration of the disease; the consequent bladder carcinoma frequently affects young people.

**Trichomoniasis**
This condition is relatively rare in males but may be acquired from an infected partner. The pyuria is relatively symptom-free, but trichomonads are found on staining the urine, or motile organisms are seen in centrifuged urine deposits. They can also be found in urethral discharge, semen or fluid massaged from the prostate.

**Urethral causes**
Urethral pyuria will be caused by any condition that causes a purulent urethritis. A profuse discharge, together with a history of recent unprotected sexual contact, is enough to provide the diagnosis, but urethritis may be secondary to cystitis as well as the converse. The symptoms of urethritis are discharge, urethral pain and occasional initial haematuria; if there is also increased frequency, suprapubic pain and bleeding throughout the stream, cystitis is also probably present. The pyuria of urethritis is usually confined to the initial sample of urine; in cystitis, the mid-stream sample will also be contaminated. Urethral calculi, foreign bodies and self-inflicted urethral trauma will also cause purulent urethritis.

**Prostatic causes**
*Acute prostatitis* presents with increased frequency, perineal and suprapubic pain, discomfort on micturition, pyuria and even acute retention. Prostatitis may arise by haematogenous spread, or it may complicate prostate biopsy, cystitis or urethritis. A rectal examination reveals a large prostate that is exquisitely tender, to the degree that touching the oedematous gland causes acute reflex contraction of the external sphincter and straightening of the hips. *Prostatic abscess* usually follows an acute urethritis that has affected the posterior urethra and caused an acute prostatitis. It may be secondary to a sexually transmitted infection, such as gonorrhoea or *Chlamydia*, and may also follow instrumentation of the urethra. The prostate is intrinsically infected subclinically, and endoscopy can trigger this infection, however carefully performed. Acute prostatitis may result in the formation of an abscess, almost always unilateral, which may discharge spontaneously into the urethra, bladder or rectum unless de-roofed by transurethral resection. Acute prostatitis presents with increased frequency of micturition, perineal and hypogastric pain, fever, rigors and difficulty with micturition. The abscess can be felt as a soft area within the tender and oedematous prostate.

Prostatic calculi are frequent, but prostatic abscesses complicating calculi are relatively unusual, as are abscesses related to genitourinary tuberculosis. Involvement of the prostate is a very late manifestation of this disease, presenting as increased frequency, perineal pain, difficulty with micturition and a sudden episode of initial haematuria.

Pyuria is invariable following prostatic surgery, whether or not covered by prophylactic antibiotics. The healing cavity of a prostatectomy, carried out transurethrally or retropubically, can take as long as 8 weeks to epithelialize, and pyuria during the whole of this period is common.

**Vesicular causes**
Seminal vesiculitis often accompanies acute prostatitis, and it often causes persistent symptoms following gonococcal or non-specific urethritis. It is a very rare complication of prostatectomy, but an abscess may develop if the openings of the vesicles are involved in the scarring process postoperatively. Tuberculous vesiculitis also occurs.
The symptoms of vesiculitis are pain in the bladder area, in the perineum and in the low back. Pyuria may be scant, but if the channels between the vesicles and urethra are free, it may be profuse. Haematospermia is not infrequent, while ascending inflammation of the vas and acute epididymitis are also often associated. The inflamed vesicle can be felt above the prostate on rectal examination. While massage can produce a bead of pus at the urethral meatus, it is difficult to distinguish this sign from the similar phenomenon encountered in acute prostatitis.

PYURIA CAUSED BY DISEASES OUTSIDE THE URINARY SYSTEM

The most common cause of pyuria is the incorrect collection of a sample, in that the urinary meatus, in the male or female, has been improperly cleaned prior to collection. In the male, retained secretions behind a phimosis can result in pyuria; an excess of physiological discharge in the female may have the same effect.

The presence and spread of inflammatory processes outside the urinary tract into the urinary passages will cause pyuria, as will the rupture of an extravesical abscess. When the symptoms suggest urinary problems (e.g. increased frequency, urgency, pain on micturition or haematuria) and are followed by the sudden appearance of a quantity of pus in the urine, there is a strong possibility of the rupture of an extra-urinary abscess into the bladder or urethra or (very rarely) into the ureter, provided that the sudden emptying of a renal abscess or pyonephrosis can be eliminated. This spontaneous discharge is often associated with a relief of the primary symptom.

The history will often provide some indication of the primary diagnosis, of which the most frequent are prostatic abscess, appendix abscess, pyosalpinx, psoas, iliac and pelvic abscess, and an abscess around a carcinoma or diverticulitis of the colon, the last of these being the most common of all.

Pyuria in acute appendicitis

If the appendix is in its usual position, the bladder is rarely affected, but if the appendix passes downwards across the pelvic brim, it is not unusual to find that the patient complains of frequency and pain on micturition when appendicitis occurs. If the appendix is severely inflamed, it may adhere to the bladder, and both pus and blood may be present in the urine. If cytoscopy is carried out, a localized area of congestion will be seen on the right lateral wall. Very occasionally, a small abscess may develop in the adhesions between the appendix and the bladder, and if this abscess discharges into the bladder, pyuria results and an enterovesical fistula is established. Diagnosis in the case of a dependent appendix is difficult; the pain is much lower in the pelvis than is usual with appendicitis, while the lower urinary tract symptoms point to a bladder disorder. The onset of the condition is gradual, however, and there is an elevation of temperature and pulse rate with right-sided abdominal rigidity. None of these is present in acute cystitis, and the possibility of an alternative acute intra-abdominal lesion must be considered. A right-sided pelvic abscess arising from a burst appendix may rupture into the bladder. The usual history of acute appendicitis is accompanied by the presence of a mass in the right iliac fossa or the pelvic space, bimanually palpable if in the latter. Pyrexia continues and is associated with rigors. If the abscess discharges into the bladder, the fever resolves and a large quantity of pus appears in the urine. Rectal examination reveals not only the tenderness of acute appendicitis but also considerable thickening relating to the thick wall of the abscess cavity.

Pyosalpinx may cause cystitis by direct spread of the inflammatory process to the bladder, and it may eventually rupture. There has usually been a history of profuse vaginal discharge associated with constant aching in the pelvic region and in the lower back; there are often frequent attacks of severe pain and malaise at variable intervals, together with an intermittent pyrexia. Periods may be profuse, frequent and more painful than usual, while vaginal examination reveals fullness or a mass in one or both vaginal fornices.

Psoas and iliac abscesses may rupture into the bladder, and the former has been known to discharge into a ureter. There is a swelling in the iliac fossa and sometimes in the inguinal region, and clinical and radiological evidence of spinal osteomyelitis, together with lateral displacement of the psoas shadow.

Diverticulitis of the pelvic colon often becomes adherent to the bladder, and if peridiverticular abscesses form, these may rupture into the bladder, causing pyuria and the formation of an enterovesical fistula. Pneumaturia – the passage of flatus per urethram – is pathognomonic, but it is surprising how rarely it occurs – the appearance of solid faecal particles in the urine is more common; when air is passed in the urine, the stream hisses or whistles. The main differential diagnosis of pneumaturia is an acute cystitis with a gas-forming organism, particularly in patients with diabetes. A colovesical fistula occurs far more frequently following the rupture of a peridiverticular abscess than by direct extension of a colonic carcinoma.
Carcinoma of the pelvic structures often involves the bladder by direct extension. This is particularly true of carcinoma of the cervix and of the rectum, but it may also occur from carcinoma of the pelvic colon, sigmoid and caecum. Spread of disease to the bladder occurs relatively late, and the symptoms of the primary condition have usually given a clear indication of the diagnosis before pyuria results. Involvement of the bladder is first shown by frequency, dysuria and urgency, while the presence of blood and pus in the urine is a late feature, representing ulceration through the whole thickness of the bladder wall. Uterovesical and vesicovaginal fistulae may result from extension of primary tumours from either of these two structures into the bladder; the pathognomonic symptom is continuous incontinence by day and night. It is hardly likely that this incontinence will need to be distinguished from that secondary to an ectopic ureter, as this will be evident from birth. Penetration of the bladder by a carcinoma of the rectum or colon will give rise to pneumaturia and the passage of pus, blood and faecal debris in the urinary stream. Occasionally, the urine flow passes in the other direction, and the urine output falls while the passage of watery stools, alternating with reasonably well-formed motions, may occur.

Tuberculosis, dysenteric ulcers of the intestines and caecal actinomycosis are rare causes of pyuria. In the last of these, the fungus, instead of infiltrating the skin and pointing in the groin externally as it usually does, extends downwards into the pelvis and opens into the bladder or rectum. The diagnosis depends on the discovery of ray fungi in the urine, and it is unlikely that they would be found unless specifically sought. Actinomycosis of the kidney is usually mistaken for tuberculosis until the fungi are discovered by microscopy.

The most common causes of the intermittent appearance of large amounts of pus in the urine are pyonephrosis, diverticulum of the bladder and vesicocolic fistula. The presence of a persistent low-grade pyuria that cannot otherwise be explained may indicate carcinoma-in-situ of the bladder or urinary tract tuberculosis.
RECTAL BLEEDING

Harold Ellis

Bleeding from the rectum is one of the most common presenting symptoms in clinical practice, and it is also the most commonly mismanaged. A classification of causes is listed in Box R.1.

The majority of patients with rectal bleeding are found to have haemorrhoids (or piles, the terms being interchangeable) as the underlying cause. Haemorrhoids are cushions of erectile tissue containing extensive arteriovenous anastomoses, and, when traumatized, arterial bleeding results (Fig. R.1). Usually, the bleeding is of a minor nature; there is light staining of the lavatory paper following defecation. Rarely, profuse bleeding leading to hypovolaemic shock can occur. The possibility of a neoplasm must always be a consideration, irrespective of the age of the patient. Although the incidence of rectal carcinoma is highest in the sixth and seventh decades of life, it is not uncommon in younger age groups. The presence of malignancy must be considered particularly when there are constitutional symptoms or there is a history of recent irregularity of bowel function. Profuse mucous discharge in association with bleeding is consistent with a villous adenoma or carcinoma, and a history of bloody diarrhoea is most consistent with a diagnosis of inflammatory bowel disease.

Approximately 80 per cent of rectal neoplasms are within range of digital examination, and a per rectum examination should be conducted in all patients with rectal bleeding. If a lesion is palpated, an assessment should be made of its mobility and fixity to surrounding tissues. A highly mobile lesion is indicative of a benign adenoma, whereas any degree of fixity is strongly suggestive of invasion and hence of malignancy. Haemorrhoids, in contrast, are not usually palpable and not tender on palpation in the absence of strangulation. Undue local tenderness suggests the presence of an underlying fissure, infection (intersphincteric abscess) or haematoma.

Sigmoidoscopy is an essential step in the exclusion of carcinoma and inflammatory bowel disease and, where there is dispute over the macroscopic appearances, biopsy is mandatory. The presence of oedema, erythema or a shallow discrete ulcer, usually confined to the anterior rectal wall, is a feature that may be indicative of the solitary rectal ulcer syndrome. This is a benign condition associated with excessive straining on defecation, and it can readily be confused with carcinoma from its sigmoidoscopic appearances.

Box R.1 Major causes of rectal bleeding

<table>
<thead>
<tr>
<th>Anal causes</th>
<th>Rectal causes</th>
<th>Colon causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Haemorrhoids</td>
<td>Angiodysplasia</td>
<td>Diverticular disease</td>
</tr>
<tr>
<td>Anal fissure</td>
<td>Ischaemia</td>
<td>Infective (e.g. dysentery)</td>
</tr>
<tr>
<td>Anal fistula</td>
<td>Infective (e.g. tuberculosis)</td>
<td>Inflammatory</td>
</tr>
<tr>
<td>Perianal haematoma</td>
<td>Inflammatory (e.g. ulcerative colitis)</td>
<td>— Ulcerative colitis</td>
</tr>
<tr>
<td>Condylomas</td>
<td>Angiodysplasia</td>
<td>Crohn’s colitis</td>
</tr>
<tr>
<td>Trauma</td>
<td>Ischaemia</td>
<td>Angiodysplasia</td>
</tr>
<tr>
<td>Malignancy</td>
<td>Neoplasia</td>
<td>Intussusception</td>
</tr>
<tr>
<td>— Squamous carcinoma</td>
<td>— Adenoma</td>
<td>Ischaemia</td>
</tr>
<tr>
<td>— Adenocarcinoma</td>
<td>— Carcinoma</td>
<td>Neoplasia</td>
</tr>
<tr>
<td>— Paget’s disease</td>
<td>— Malignant melanoma</td>
<td>— Adenoma</td>
</tr>
<tr>
<td>— Malignant melanoma</td>
<td></td>
<td>— Carcinoma</td>
</tr>
<tr>
<td>— Bowen’s disease</td>
<td></td>
<td>General causes</td>
</tr>
<tr>
<td>— Basal cell carcinoma</td>
<td></td>
<td>Clotting deficiencies</td>
</tr>
<tr>
<td>Rectal caused</td>
<td></td>
<td>Anticoagulants</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Uraemia</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Trauma</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Neoplasia</td>
</tr>
<tr>
<td></td>
<td></td>
<td>— Adenoma</td>
</tr>
<tr>
<td></td>
<td></td>
<td>— Carcinoma</td>
</tr>
</tbody>
</table>

Figure R.1 Third-degree haemorrhoids, or piles. The patient is lying on the operating table in the lithotomy position awaiting haemorrhoidectomy.
The diagnosis of haemorrhoids largely rests on the appearances at proctoscopy. Most commonly, the right anterior haemorrhoid is noted to be enlarged and congested, and is the putative cause of bleeding since it is rare to see the active bleeding source at the time of the examination.

Patients with a history of blood mixed in with the stool, or where there is a major loss of altered or of venous blood, may require more detailed investigation, which will include barium enema and colonoscopy. If angiodyplasia is to be excluded, arteriography may be necessary. Finally, rectal bleeding may be readily confused with bleeding from the upper gastrointestinal tract and small intestine (see MELAENA, p. 399).

RECTAL DISCHARGE

Harold Ellis

The causes of rectal discharge are classified and listed in Box R.2.

Secretion from sweat glands in the perianal area and from the anal glands is a common and normal phenomenon that rarely gives rise to significant problems. Profuse mucous secretion, however, often causes considerable discomfort and pruritus ani as a consequence of inflammation of the perianal skin. Such secretion is commonly observed with haemorrhoids, particularly where there is a combination of prolapse with a weak internal anal sphincter. More serious pelvic floor disorders (e.g. complete prolapse or solitary rectal ulcer syndrome) may be responsible for profuse and sometimes blood-stained mucous secretion, which can be incapacitating. Inflammation of the rectal mucosa from ulcerative colitis or Crohn’s disease of the rectum may similarly produce a mucous discharge that is usually blood-stained and accompanied by diarrhoea. The existence of neoplasia should always be suspected, since copious secretion is a particular feature of villous adenomas of the rectum in which the potassium loss may be sufficient to induce hypokalaemia. Carcinoma may also be a cause of mucous secretion, although bleeding is usually a more prominent feature and there may be constitutional symptoms.

A purulent discharge is usually caused by anal and perianal sepsis. On inspection of the perineum, a small opening discharging pus to the side of the anus is highly suggestive of a fistula (Fig. R.2). The diagnosis can be confirmed by palpation and observation at proctoscopy of the internal opening. Ulcerating and purulent perianal lesions should raise the possibility of Crohn’s disease (Fig. R.3), anal tuberculosis or sexually transmitted disease (e.g. syphilis, gonorrhoea or HIV infection). Where there is doubt, bacteriological examination of the pus and histological examination of the biopsy from the perianal skin should be performed. Anal neoplasms and condylomas can be responsible for an offensive purulent discharge; the diagnosis is apparent on inspection, but biopsy is always mandatory even if simple condyloma is diagnosed since malignant development can occur with this lesion.

**Box R.2**  Classification of major causes of rectal discharge

<table>
<thead>
<tr>
<th>Discharge of mucus</th>
<th>Discharge of pus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Haemorrhoids</td>
<td>Anal fistula</td>
</tr>
<tr>
<td>Rectal prolapse</td>
<td>Perianal Crohn’s disease</td>
</tr>
<tr>
<td>Solitary rectal ulcer syndrome</td>
<td></td>
</tr>
<tr>
<td>Villous adenoma</td>
<td>Anal tuberculosis</td>
</tr>
<tr>
<td>Carcinoma of the rectum</td>
<td>Anal neoplasms</td>
</tr>
<tr>
<td>Proctitis</td>
<td>Anal fissure</td>
</tr>
<tr>
<td></td>
<td>Syphilis</td>
</tr>
<tr>
<td></td>
<td>Gonorrhoea</td>
</tr>
<tr>
<td></td>
<td>Condyloma accuminata</td>
</tr>
</tbody>
</table>

**Figure R.2** Fistula-in-ano. The patient is on the operating table in the lithotomy position. Under anaesthetic, a probe has been passed through the external opening of the fistula at the 5 o’clock position and can be seen to emerge through the internal opening in the midline.
Every medical practitioner should be aware of the importance of conducting a digital examination of the anal canal and rectum in all patients with anorectal symptoms, since the majority of rectal neoplasms are well within reach of the examining finger. The relevance of performing a rectal examination as part of a general examination in a patient without rectal symptoms is less clear. Since rectal cancer is a common malignancy in patients aged over 60 years, a strong argument can be made that all patients in this age group should undergo rectal examination as part of any general physical examination. Clearly, if there are urinary symptoms, a digital assessment of the prostate is highly relevant, and similarly digital examination of the rectum (and, where relevant, vaginal examination) may be valuable in patients with pelvic or perineal symptoms.

Digital examination of the rectum should be conducted, where possible, with the patient lying in the left lateral position, and should not be attempted until a full inspection of the perineum has been conducted to exclude fissure or other pathology which might give rise to severe pain on palpation. Initially, a digital assessment is made of anal sphincter tone, which may be increased in the presence of fissure and decreased in functional disorders such as anorectal incontinence.

Each quadrant of the anus and rectum should be examined sequentially. Within the anus, lesions may extend caudally from the rectum, and vice versa. In the normal state, haemorrhoids are not palpable, and no specific structure is palpated until the examining finger reaches the rectum. In women, the cervix frequently projects into the anterior rectal wall and is readily palpable; this is frequently mistaken by an inexperienced clinician for a rectal neoplasm, as may a vaginal tampon! In men, the prostate is easily palpable anteriorly. Laterally, the ischial spines may be palpated, and this may be of value in the location of the pudendal nerves (to provide a pudendal nerve blockade). Posteriorly, the shelf created by the levator ani, and in thin subjects, the bony coccyx may be palpable.

A classification of major causes of rectal masses is provided in Box R.3.

<table>
<thead>
<tr>
<th>Intrinsic causes</th>
<th>Rectal neoplasia</th>
</tr>
</thead>
<tbody>
<tr>
<td>Benign</td>
<td>Polyp*</td>
</tr>
<tr>
<td>Villous adenoma*</td>
<td></td>
</tr>
<tr>
<td>Leiomyoma</td>
<td></td>
</tr>
<tr>
<td>Malignant</td>
<td>Carcinoma*</td>
</tr>
<tr>
<td>Carcinosarcoma</td>
<td></td>
</tr>
<tr>
<td>Leiomyosarcoma</td>
<td></td>
</tr>
<tr>
<td>Lymphoma</td>
<td></td>
</tr>
<tr>
<td>Melanoma</td>
<td></td>
</tr>
</tbody>
</table>

| Anal neoplasia |
| Benign | Condylomas* |
| Polyps* |
| Malignant | Adenocarcinoma* |
| Squamous carcinoma* |
| Melanoma |
| Bowen's disease |
| Paget's disease |
| Basal cell carcinoma |

| Infection | Lymphogranuloma |
| Tuberculosis |

| Extrinsic causes | Benign prostatic hypertrophy* |
| Infection | Pelvic abscess |
| Anal fistula with suprarectal extension |

| Tumour | Secondary spread to pelvis/pouch of Douglas |
| Carcinoma of the prostate* |

| Gynaecological causes |
| Carcinoma of the body of the uterus* |
| Carcinoma of the cervix* |
| Carcinoma of the ovary* |
| Fibroid uterus* |
| Ovarian cyst* |
| Pyosalpinx |
| Ectopic gestation |
| Endometriosis |

| Presacral (retrorectal) causes |
| Congenital | Epidermoid cyst |
| Teratoma/carcinoma |
| Meningocele |
| Chordoma |

| Causes arising from bone/cartilage | Osteogenic sarcoma |
| Exing's sarcoma |
| Osteochondroma |
| Myeloma |
| Giant-cell tumour |

| Neurological causes | Neurilemmoma |
| Ependymoma |
| Neuroblastoma |

| Miscellaneous | Lipoma |
| Haemangioma |

*Common causes are indicated by asterisks
INTRINSIC LESIONS

If a mass is perceived on digital examination, it is not always possible to decide on palpation alone if the lesion is intrinsic or extrinsic. The differentiation may only be possible after sigmoidoscopy and histological examination. The consistency and mobility may closely relate to the diagnosis. Hence, a benign villous adenoma (Fig. R.4) will feel soft, fleshy and highly mobile. In contrast, a carcinoma (Fig. R.5) may feel hard, with obvious fixation of the mucosal lesion to the underlying muscle or perirectal fat. Sometimes, nearby extrarectal lymph nodes containing secondary tumour deposits may be palpable.

A circumferential stenosis of the rectum may be seen following trauma (e.g. previous surgery) or be a complication of (i) infection (e.g. lymphogranuloma); (ii) inflammation (e.g. ulcerative colitis); or (iii) ischaemia. Digital examination in these circumstances is usually accompanied by marked tenderness and pain. Anal neoplasms may extend from the anus and upwards into the rectum, in which case there may be a marked stenosis, and the examination will cause pain.

EXTRINSIC LESIONS

Rectal examination is a simple clinical means of diagnosing the presence of pelvic pus or pelvic tumour. A tender mass in the presence of oedematous rectal mucosa suggests a collection of pus, whereas a hard, fixed extrinsic mass in which there is no mobility of the rectal wall or uterus would be strong evidence in favour of pelvic malignancy. Infection may arise secondary to gynaecological or intestinal sepsis, but it may be secondary to an anal fistula where pus has tracked superiorly to create a collection above the levator musculature. Such fistulas are important to recognize since their treatment is complex.

Benign enlargement of the prostate may give rise to symmetrical hypertrophy of the gland in which the midline sulcus is preserved. Malignant enlargement gives rise to a mass that is denser and asymmetrical, and the midline sulcus is invaded.

A mass that is clearly situated posterior to the rectum arises in the potential space ventral to the sacrum and coccyx bounded distally by the levator ani, and proximally by the pelvic peritoneal reflection. The important primary distinction is whether the lesion is solid or cystic; this may require ultrasound examination for confirmation. The majority of solid lesions are chordomas, and a cystic lesion is usually one of the following: (i) epidermoid cyst; (ii) mucus-secreting cyst; (iii) teratoma; (iv) teratocarcinoma; or (v) meningocoele. Neurogenic and osseous tumours are rare in this region. Clinically, presacral masses usually present with a history of low back pain radiating to the rectum and buttocks. Pressure on the bladder may lead to urinary retention, and constipation is a frequent symptom. On sacral radiographs, presacral lesions may show up as an area of calcification with rarefaction of the sacrum; if there is bony destruction,
malignancy should always be suspected. A barium enema should always be performed to exclude colonic communication with the mass; and similarly, communication with the subarachnoid space should be excluded by CT or magnetic resonance imaging.

**RECTAL TENESMUS**

Harold Ellis

Rectal tenesmus is a non-specific term employed to describe a state in which there is either difficulty with or repeated, painful and sometimes futile defecation. A similar condition has been described affecting micturition and is referred to as urinary tenesmus (or strangury). The repeated defecation is often accompanied by the passage of mucus and/or blood, and this collection of symptoms should be distinguished from the symptom of diarrhoea. In the latter, there is either a complaint of stool of loose consistency or there is increase in frequency but, in contrast to patients with tenesmus, defecation is usually productive of stool.

As with all rectal symptoms, there may be a sinister underlying cause, and a full clinical assessment – which will include digital examination of the anus/rectum and sigmoidoscopy – is essential.

**INFLAMMATORY AND INFECTIVE CAUSES (PROCTITIS)**

Proctitis (see RECTAL ULCERATION, below) may be either inflammatory (e.g. ulcerative colitis or Crohn’s disease) or infective in origin. The tenesmus may be associated with constitutional symptoms (e.g. malaise, weight loss and anorexia) and with severe diarrhoea. In patients with ulcerative colitis, the bleeding may be substantial, leading to severe anaemia. The diagnosis is readily made by sigmoidoscopy, biopsy and, where relevant, stool culture. Less commonly, perianal sepsis (e.g. fistula) can cause tenesmus. The diagnosis is suggested either by the presence of extreme tenderness on digital examination of the anus or by the presence of a sinus/fistula opening in the perianal region.

**NEOPLASTIC CAUSES**

Benign (e.g. villous adenoma) or malignant lesions of the rectum or anus frequently cause tenesmus. An extensive villous adenoma of the rectum is notorious as a cause of excessive secretion of rectal mucus, which is sufficiently rich in potassium to lead to hypokalaemia. Adenocarcinoma of the rectum may similarly be responsible for the secretion of mucus, but to a lesser degree. Rectal bleeding is a more pronounced feature in the history; the bleeding may be bright or dark red, may be mixed in with the stool, and accompanies defecation. In the case of advanced malignancy, the rectal symptoms may be accompanied by constitutional symptoms, such as weight loss.

Squamous carcinoma of the anal margin should be recognizable on simple inspection of the anal verge, and examination of the inguinal region may reveal lymphadenopathy in the presence of metastatic spread to the regional nodes.

**MECHANICAL CAUSES**

Tenesmus occasionally results from a poorly understood condition in which the pelvic floor and external anal sphincter musculature fail to relax or may actively contract during attempted defecation. In normal circumstances, these muscles relax reflexly to enable the easy passage of the faecal bolus through the anal canal. This condition is usually diagnosed only either by conventional electromyography or by defecography (radiographic imaging of the rectum during defecation after barium installation), and may be associated with a solitary rectal ulcer as seen on sigmoidoscopy (see RECTAL ULCERATION, below). The cause is usually not known, but pelvic floor ‘spasticity’ is occasionally identified in patients with multiple sclerosis, and the symptom of tenesmus may be the first symptom noted by patients with demyelinating diseases.

Minor anorectal disorders, particularly in an acute presentation (e.g. perianal thrombosis), may cause tenesmus, since the lesion within the anus may cause stimulation of the anal sensory receptors at and below the dentate line, which gives rise to a false impression that there is faecal matter present within the anus and lower rectum.

**RECTAL ULCERATION**

Harold Ellis

Normally, a diagnosis of rectal ulceration will be made from the macroscopic appearances of the rectum either at sigmoidoscopy or radiologically at barium enema. Only in certain circumstances (see below) will an ulcer be palpable.

The major causes of rectal ulceration are listed in Box R.4.

The clinical distinction between inflammatory bowel disease and infection can rarely be made on macroscopic appearances alone. Hence, ulcerative colitis and Shigella infection may both give rise to: (i) shallow ulceration; (ii) granular appearances; (iii) haemorrhagic friable mucosa; and (iv) oedematous mucosa in the rectum. The presence of pseudopolyps

**Box R.4**

**CAUSES**

**RECTAL TUMOURS**

**NEOPLASTIC CAUSES**

Benign (e.g. villous adenoma) or malignant lesions of the rectum or anus frequently cause tenesmus. An extensive villous adenoma of the rectum is notorious as a cause of excessive secretion of rectal mucus, which is sufficiently rich in potassium to lead to hypokalaemia. Adenocarcinoma of the rectum may similarly be responsible for the secretion of mucus, but to a lesser degree. Rectal bleeding is a more pronounced feature in the history; the bleeding may be bright or dark red, may be mixed in with the stool, and accompanies defecation. In the case of advanced malignancy, the rectal symptoms may be accompanied by constitutional symptoms, such as weight loss.

Squamous carcinoma of the anal margin should be recognizable on simple inspection of the anal verge, and examination of the inguinal region may reveal lymphadenopathy in the presence of metastatic spread to the regional nodes.

**MECHANICAL CAUSES**

Tenesmus occasionally results from a poorly understood condition in which the pelvic floor and external anal sphincter musculature fail to relax or may actively contract during attempted defecation. In normal circumstances, these muscles relax reflexly to enable the easy passage of the faecal bolus through the anal canal. This condition is usually diagnosed only either by conventional electromyography or by defecography (radiographic imaging of the rectum during defecation after barium installation), and may be associated with a solitary rectal ulcer as seen on sigmoidoscopy (see RECTAL ULCERATION, below). The cause is usually not known, but pelvic floor ‘spasticity’ is occasionally identified in patients with multiple sclerosis, and the symptom of tenesmus may be the first symptom noted by patients with demyelinating diseases.

Minor anorectal disorders, particularly in an acute presentation (e.g. perianal thrombosis), may cause tenesmus, since the lesion within the anus may cause stimulation of the anal sensory receptors at and below the dentate line, which gives rise to a false impression that there is faecal matter present within the anus and lower rectum.

**RECTAL ULCERATION**

Harold Ellis

Normally, a diagnosis of rectal ulceration will be made from the macroscopic appearances of the rectum either at sigmoidoscopy or radiologically at barium enema. Only in certain circumstances (see below) will an ulcer be palpable.

The major causes of rectal ulceration are listed in Box R.4.

The clinical distinction between inflammatory bowel disease and infection can rarely be made on macroscopic appearances alone. Hence, ulcerative colitis and Shigella infection may both give rise to: (i) shallow ulceration; (ii) granular appearances; (iii) haemorrhagic friable mucosa; and (iv) oedematous mucosa in the rectum. The presence of pseudopolyps
If rare organisms are cultured, such as the protozoan *Legionella*, only the more important are listed in Box R.4. The list of infective agents that can give rise to a proctitis is legion; only the more important are listed in Box R.4. If rare organisms are cultivated, such as the protozoan cryptosporidium or viruses (e.g. cytomegalovirus or *herpes simplex*), the possibility of immune deficiency (e.g. HIV infection or leukaemia) should always be considered.

Ischaemia rarely affects the rectum; when it does, however, it is usually prevalent in older age groups, is of sudden onset, and is associated with profuse bleeding and abdominal pain. The diagnosis is confirmed by barium enema in which the characteristic appearances of ‘thumb-printing’ are observed principally in the descending and sigmoid colon.

The ulcer of the solitary rectal ulcer syndrome and the ulcerating lesion associated with some rectal carcinomas are often both readily palpable on rectal examination. Both may feel indurated with fixity to extrarectal tissues, and the sigmoidoscopy appearances can be identical. Adequate biopsy is essential to enable the diagnosis to be made since the presence of carcinoma will require radical surgical measures. Ulceration may rarely be traumatic in origin, either as a result of self-mutilation or because digitally assisted evacuation is the only means by which the voiding of rectal contents can be achieved.

### REFLEXES, ABNORMALITIES OF

**David Werring & Mark Kinirons**

A reflex, first conceived by Descartes, is the most basic form of involuntary response to a stimulus. The anatomical pathway (reflex arc) consists of:

- (i) a receptor organ;
- (ii) an afferent pathway to the brain or spinal cord;
- (iii) usually interneurones in the central nervous system; and
- (iv) an efferent path that leaves via the lower motor neurones to reach a receptor organ, often a muscle. The stimulus may be touch, pinprick or sudden stretching of a muscle. The response may be not only muscle contraction but also altered muscle tone or glandular secretion.

A reflex depends upon the integrity of its reflex arc. Lesions of the pathway abolish the reflex. This could be due to damage to the peripheral nerve or central pathways. Reflexes are profoundly modulated by descending pathways, including the corticospinal (pyramidal) and parapyramidal tracts (e.g. reticulospinal tract, vestibulospinal tract and rubrospinal tract). Damage to the pyramidal and parapyramidal tracts causes exaggerated reflexes below the level of the lesion; thus, hyperreflexia constitutes one component of the upper motor neurone syndrome (together with hypertonia, associated reactions, spastic dystonia, co-contraction, etc.). Anxiety and pain, and systemic illnesses including hyperthyroidism, may also cause...

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<table>
<thead>
<tr>
<th>Box R.4 Major causes of rectal ulceration</th>
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<tbody>
<tr>
<td>- Inflammatory</td>
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<tr>
<td>- Ulcerative colitis</td>
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<tr>
<td>- Crohn's disease</td>
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<tr>
<td>- Post radiation</td>
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<tr>
<td>- Infective</td>
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<tr>
<td>- Shigella</td>
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<td>- Salmonella</td>
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<td>- Campylobacter</td>
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<td>- Tuberculous</td>
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<tr>
<td>- Gonococcal</td>
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<tr>
<td>- Amoebias</td>
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<tr>
<td>- Pseudomembranous colitis (Clostridium difficile)</td>
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<tr>
<td>- Lymphogranuloma</td>
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<tr>
<td>- Schistosoma</td>
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<td>- Syphilis</td>
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<tr>
<td>- <em>Herpes simplex</em></td>
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<tr>
<td>- Enterovirus</td>
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<tr>
<td>- Cytomegalovirus</td>
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<tr>
<td>- HIV infection</td>
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<tr>
<td>- Solitary rectal ulcer syndrome</td>
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<tr>
<td>- Trauma</td>
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<tr>
<td>- Malignant ulcer</td>
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<tr>
<td>- Carcinoma</td>
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<tr>
<td>- Leukaemia</td>
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<tr>
<td>- Ischaemia</td>
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hyperreflexia. In conditions causing hypotonia (e.g. cerebellar disease), the tendon reflexes are depressed. Some otherwise neurologically normal individuals may have tendon reflexes that cannot be elicited, and loss of these reflexes is (together with tonic pupils) a part of the Holmes–Adie syndrome. Poor relaxation does not allow successful tendon reflex testing. The main reflexes are listed in Box R.5; of these, a comparatively small number are of diagnostic value in clinical neurology. Only the clinically useful reflexes will be discussed here. The pupillary and important brainstem reflexes are considered elsewhere (see PUPILS, ABNORMALITIES OF; p. 551; and NYSTAGMUS, p. 466).

THE TENDON REFLEXES

The tendon reflex is a reflection of the myotactic reflex: a reflex contraction of a muscle (or a part of a muscle) in response to rapid stretch, as caused by tapping with a tendon hammer. It is a monosynaptic reflex; the reflex pathway ascends from the muscle through the posterior roots, passes forward to the anterior horn cells of the same segment, and descends to the muscle or muscles concerned via the motor nerve. Anatomical factors limit the practical application of this test to the muscles listed in Table R.1.

Exaggerated tendon reflexes

The tendon reflexes vary widely in different individuals, and in a given individual at different times. The muscles involved must be relaxed during testing, and patients should be calm and generally relaxed, otherwise the reflexes will be misleadingly brisk. Localized exaggeration of tendon reflexes is a valuable sign of damage to the corticospinal (pyramidal) pathways above the reflex. Damage to the corticospinal tracts also reduces superficial cutaneous reflexes (abdominal and cremasteric). In addition to pathological exaggeration, reflex ‘spread’ to muscles innervated by other segments is another sign of pathology above the lesion. The supinator jerk may spread to the finger flexors in cases of cervical myelopathy. In the arms, ‘inversion’ of the reflexes may be of localizing value. For example, a lesion at C5–C6 involving the cervical nerve roots that is also compressing the corticospinal tracts (e.g. a local intervertebral disc protrusion at this level) will cause a reduced biceps jerk on tapping the tendon but contraction of the triceps (which is the ‘inverted’ reflex) instead, as well as exaggerated reflexes below this level. Similarly, the supinator may be lost, with finger flexion at the segment below instead. Hoffman’s reflex may also be associated with hyperreflexia due to cervical lesions above C7: it is elicited by flexing the distal interphalangeal joint of the middle finger and then flicking it down further such that the finger springs back to the normal position. A positive Hoffman reflex is brisk thumb contraction.

Depression or loss of tendon reflexes

This may occur as a transient event in ‘spinal shock’ after severe, sudden traumatic or vascular injury to the spinal cord. General cerebral depression (e.g. deep anaesthesia or traumatic brain injury), systemic illness (e.g. severe general infections, diabetic coma or uraemia) or drugs (e.g. hypnotics or sedating anticonvulsants) may also depress the tendon reflexes. A few otherwise neurologically normal people have absent tendon reflexes. It is useful to compare the same reflex on both sides of the body, and in the upper

## Table R.1 Common tendon reflexes and their segmental values

<table>
<thead>
<tr>
<th>Reflex</th>
<th>How to elicit it</th>
<th>Response</th>
<th>Segmental value</th>
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<tbody>
<tr>
<td>Biceps</td>
<td>Tab biceps tendon</td>
<td>Flexion of elbow</td>
<td>C5–C6</td>
</tr>
<tr>
<td>Triceps</td>
<td>Tab triceps tendon</td>
<td>Extension of elbow</td>
<td>C6–C7</td>
</tr>
<tr>
<td>Supinator</td>
<td>Tab brachioradialis tendon (distal end of the radius)</td>
<td>Flexion elbow</td>
<td>C5–C6</td>
</tr>
<tr>
<td>Finger jerk</td>
<td>Tab palmar surface of semiflexed, relaxed fingers</td>
<td>Flexion of fingers (and thumb)</td>
<td>C7–C8</td>
</tr>
<tr>
<td>Knee jerk</td>
<td>Tab quadriceps tendon below</td>
<td>Extension of knee</td>
<td>L2–L4</td>
</tr>
<tr>
<td>Ankle jerk</td>
<td>Tab tendo calcaneus (Achilles)</td>
<td>Plantarflexion of ankle</td>
<td>S1–S2</td>
</tr>
</tbody>
</table>
and lower limbs, before making a judgement. When no reflex can be obtained, reinforcement is used by the forcible contraction of muscles remote from the tested segment (e.g. forced hand-clenching to reinforce the knee and ankle jerks, or clenching the teeth to reinforce the upper limb tendon jerks). Segmental loss of tendon reflexes is a very useful localizing sign in neurology. Any part of the reflex arc (muscle, peripheral nerve or spinal cord) may be interrupted, leading to loss of the appropriate tendon reflex.

**DISEASE OF MUSCLE**
Depression or loss of reflexes occurs as a late event in any form of myopathy, but it is usually widespread, adding little additional localizing value to the other signs of myopathy, including muscle wasting.

**DISEASE OF PERIPHERAL NERVES**
Damage to either the motor or sensory peripheral nerves may lead to a loss of tendon reflexes. This may occur where the damage is localized to one or more individual peripheral nerves (mononeuritis multiplex), or where the damage occurs as part of a generalized peripheral neuropathy. Involvement of a single nerve is manifest as motor and sensory signs conforming to a single nerve territory. The differential diagnosis of mononeuritis multiplex is listed in Box R.6. Mononeuropathy, usually isolated to one nerve, can also result from entrapment (e.g. carpal tunnel syndrome due to median nerve compression at the wrist, or cubital tunnel syndrome due to ulnar nerve compression at the elbow).

There are many causes of generalized polyneuropathy, and a detailed discussion is beyond the scope of this book. The more important ones are listed in Box R.7. In clinical practice, Guillain–Barré syndrome is the most important cause of an acute sensorimotor polyneuropathy, while diabetes, alcohol and chronic inflammatory demyelinating polyradiculopathy are common causes of a chronic sensorimotor polyneuropathy. Neurophysiological characterization of the neuropathy as predominantly demyelinating (with slowed conduction) or axonal (with reduced sensor and motor action potential amplitudes) can be helpful in narrowing the differential diagnosis. Demyelinating neuropathies include Guillain–Barré syndrome, chronic inflammatory demyelinating polyneuropathy, paraproteinaemic neuropathy and Charcot–Marie–Tooth disease (type 1).

**DISEASE OF THE MOTOR OR SENSORY SPINAL ROOTS**
Damage to either the afferent or efferent part of the spinal reflex as a result of root damage is a common cause of loss of a localized tendon reflex. Common causes are cervical and lumbar spondylosis, with or without disc herniation, intraspinal tumours, and brachial or lumbar plexopathy (e.g. from neoplastic

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**Box R.6 Causes of mononeuritis multiplex**

- Diabetes
- Ischaemic neuropathy
- Vasculitis
- Polyarteritis nodosa
- Churg–Strauss syndrome (with asthma/eosinophilia)
- Wegener’s granulomatosis
- Essential mixed cryoglobulinaemia
- Rheumatoid arthritis
- Lupus
- Brachial neuritis
- Hepatitis C (with or without cryoglobulinaemia)
- HIV
- Sarcoidosis
- Lyme disease
- Sjögren–sicca syndrome (dorsal root ganglionopathy)
- Chronic ataxic neuropathy (idiopathic sensory ganglionopathy)
- Migrant polyneuritis of Wartenburg
- Coeliac disease

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**Box R.7 Causes of generalized polyneuropathy**

**Acute motor paralysis**

- Guillain–Barré syndrome
- Diphtheria
- Porphyria

**Subacute sensorimotor paralysis**

- Vitamin B12 deficiency
- Beri-beri
- Alcohol
- Drugs (e.g. isoniazid, chemotherapy [vincristine, cisplatin, etc.], phenytoin, chloramphenicol)
- Heavy metal poisoning (e.g. arsenic, mercury, lead, thallium)

**Chronic sensorimotor neuropathies**

- Associated with neoplasia: carcinoma, lymphoma, myeloma, paraproteinaemias, others
- Chronic inflammatory demyelinating polyneuropathy
- Amyloid
- Diabetes
- Leprosy

**Inherited neuropathies**

- Hereditary motor–sensory neuropathy (Charcot–Marie–Tooth)
- Hypertrophic (Djerine–Sottas disease)
- Hereditary liability to pressure palsies
- Metabolic (Refsum’s disease, leucodystrophies, abetalipoproteinaemia)
- Mitochondrial diseases
- Riley–Day syndrome

**Spinocerebellar degenerations**
infiltration or radiation). Tabes dorsalis produces areflexia due to damage to the posterior nerve roots.

**DISEASE OF THE SPINAL CORD**

Damage within the spinal cord, either within the dorsal root entry zone, between the dorsal root entry zone and the anterior horn cell, or in the anterior horn cell itself, may produce loss of the tendon reflex. This may occur in intramedullary lesions, such as syringomyelia or intramedullary tumours. A syrinx classically gives dissociated sensory loss (reduced pain and temperature sensation, but preserved joint position and light touch) with reduced reflexes at the involved segmental level, commonly C5–C6. In motor neurone disease, there is usually damage to both upper and lower motor neurones, so that the reflexes are pathologically brisk in wasted (denervated) muscles.

**REGURGITATION**

Simon Anderson

In regurgitation, the patient is aware of food that is passed from the oesophagus into the mouth. There is therefore relaxation of the upper oesophageal sphincter to allow the contents of the oesophagus to enter the mouth. It is important to distinguish between regurgitation, gastro-oesophageal reflux and vomiting. In vomiting, the gastric contents pass through open lower and upper oesophageal sphincters as a consequence of forceful contractions of the abdominal wall and the stomach muscles. Gastro-oesophageal reflux (GOR) is the passive passage of gastric acid and bile into the oesophagus. Regurgitation can occur in severe GOR (the terms are often used synonymously), but this is usually indicative of a motility disorder of the oesophagus or a behavioural disorder (rumination syndrome).

While regurgitation can be regarded as a classical symptom of oesophageal disorder, it is not necessarily so, and it is commonly seen in infants, where regurgitation can be a common and normal phenomenon and is related to the passage of gastric contents into the child’s mouth.

Patients who complain of regurgitation will often indicate that there is a postural element, with the symptom being most marked by change of position, particularly when bending forward and often on physical exercise. The symptom occurs classically in achalasia. In this motor disorder of the oesophagus, there is a reduction in the number of ganglion cells that innervate the musculature of the lower oesophageal sphincter. A weakness of the oesophageal body typically ensues. The patient is usually between 20 and 40 years of age, and classically describes dysphagia, painful swallowing and regurgitation. The regurgitated food is classically bland-tasting and contains no stomach acid. When this occurs at night-time, aspiration pneumonia can occur. Halitosis and chest pain are other symptoms. The diagnosis is made by oesophageal manometry studies. The resting pressure in the lower oesophageal sphincter is usually elevated and fails to fall, as is normal with swallowing. A poorly contracting, dilated oesophagus will be seen on barium swallow, with the non-relaxing lower oesophageal sphincter giving the appearance of a ‘beak’ on the X-ray (Fig. R.6). Upper endoscopy should be performed to exclude a tumour of the lower oesophagus (which can mimic achalasia). Follow-up endoscopies are needed as surveillance for the development of squamous carcinoma of the oesophagus, which is a recognized complication of long-standing achalasia.

A picture very similar to achalasia is produced by Chagas disease due to Trypanosoma cruzi, which is encountered in South America. The disease is characterized by a cardiomyopathy and oesophageal dysmotility which can be mild or severe, resulting in a mega-oesophagus and sometimes a megacolon. It is the cardiac complications of the disease that usually bring the patient to medical attention, but occasionally

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**Figure R.6** Barium swallow showing a dilated oesophagus with the typical rat’s tail appearance (arrowed) due to achalasia.
the oesophageal symptoms may be dominant, and they are then identical to those of classic achalasia. Other motor disorders that may cause regurgitation and dysphagia include the collagen vascular disorders such as scleroderma, diabetes mellitus and alcoholic neuropathy. In these conditions, reflux or regurgitation is not a prominent feature, dysphagia and heartburn being more troublesome.

Gastro-oesophageal reflux can result in reflux oesophagitis – inflammation of the lower oesophagus and potentially the development of Barrett's mucosa (see HEARTBURN, p. 276). The two terms are not synonymous, however, because reflux into the oesophagus from the stomach is not necessarily associated with macroscopic inflammation. Even significant acid reflux may not be evident at endoscopy. Reflux is associated with a reduced tone in the lower oesophageal sphincter, either in the resting state or transiently (commonly after meals). In addition to this, secondary peristaltic clearing of often normal amounts of gastric refluxate from the oesophagus is inadequate. In other words, instead of refluxed material being promptly cleared back into the stomach, it remains for a longer period than normal in the oesophagus, thereby causing symptoms and possibly inflammation. A reduced lower oesophageal sphincter pressure has been seen following the ingestion of fat and alcohol. Smoking also tends to cause relaxation of the oesophageal sphincter, as do some drugs such as morphine, pethidine, oestrogens and diazepam. Carminatives such as coffee will also cause a temporary relaxation of the sphincter.

A sliding hiatus hernia, if significant size, is not necessarily associated with reflux or regurgitation. Some patients may have a relatively insensitive oesophagus and not perceive refluxed acid in the oesophagus. A low gastric acidity (typically in the elderly) or associated with chronic helicobacter pylori infection may mean less acid passes into the lower oesophagus.

The functioning of the oesophagus and upper and lower oesophageal sphincters can be accurately measured by oesophageal manometry (by the placement of a nasal-oesophageal catheter) and 24-hour pH study. A barium swallow can also add information.

It is well worth stressing that, in many patients who have reflux as manifested by heartburn, or occasionally by regurgitation of food into the mouth, no clear aetiological factor can be determined. Tests of oesophageal motor function and 24-hour monitoring of oesophageal pH may not demonstrate any evidence of abnormality. These are important considerations in deciding whether or not a patient with regurgitation and a demonstrable hiatus hernia should be subjected to surgery.

RUB, PERICARDIAL

Gerry Carr-White

Inflammation of the pericardium may cause the visceral and parietal layers of the pericardium to produce audible friction as they rub against each other. A pericardial rub is therefore pathognomonic of acute pericarditis. A rub has a characteristic ‘creaking’ or ‘leathery’ character, but this quality does not always clearly differentiate it from a cardiac murmur. A typical rub is virtually continuous throughout the cardiac cycle, but three components have been identified: systolic, early diastolic and, if the patient is in sinus rhythm, late diastolic. If all three components are heard, there is little doubt about the diagnosis, but the to-and-fro cadence of the systolic and early diastolic components can sometimes be confused with the murmurs of aortic stenosis and regurgitation. If the rub is heard in systole only, as is often the case when the pericarditis is resolving, it may easily be confused with a murmur.

A rub may be heard anywhere over the precordium. It may be markedly influenced by respiration and posture (increasing on lying down and reducing on sitting up), although the changes are not predictable and, to be sure of not missing a rub, it is necessary to auscultate in all phases of deep respiration and with the patient in several postures. Sometimes it may be possible to increase the intensity of a rub by increasing the pressure with which the stethoscope is applied to the chest. In practice, there is rarely much difficulty in identifying a pericardial rub, but there may occasionally be confusion with the ‘scratchy’ murmur of Epstein's anomaly or the late systolic murmur of mitral valve prolapse. When doubt remains, the passage of time will usually resolve the issue as pericardial rubs are evanescent and usually disappear within a few days.

The causes of pericarditis are discussed under CHEST, PAIN IN (p. 88).
SALIVARY GLANDS, PAIN IN

Harold Ellis

Pain in one or other of the major salivary glands is associated with enlargement of the affected organ itself (see SALIVARY GLANDS, SWELLING OF, p. 596). For practical purposes, this symptom is confined to the parotid and submandibular salivary glands. Painful enlargement of the sublingual gland is rare; it is occasionally seen as a manifestation of mumps, together with painful enlargement of the other glands, and in the unusual condition of an advanced carcinoma of the gland itself or invasion from adjacent structures. This gland will not be considered further.

The painful salivary swellings may be classified as follows:

- **Parotid gland**
  - Mumps (epidemic parotitis)
  - Acute bacterial suppurative parotitis – postoperative, dehydration, following radiotherapy
  - Parotitis association with duct obstruction (calculus, trauma)
  - Carcinoma (primary or spread from another focus)

- **Submandibular gland**
  - Mumps (rare)
  - Inflammation associated with duct obstruction
  - Carcinoma

**MUMPS**

Mumps (epidemic parotitis) is a viral disease that is transmitted by droplet infection and has an incubation period of 17–21 days. It is the most common cause of a painful parotid swelling.

Children are most often affected. There is usually prodromal fever with malaise. Only one gland may be involved, or both may be affected simultaneously, or one gland may become enlarged after the other. The swelling progresses for several days with a marked tenderness of the gland and thickening of the overlying skin. There is characteristic uplifting of the lobe of the ear and stiffness of the jaw.

Rarely, the submandibular glands may also become swollen, and this may also unusually implicate the sublingual glands.

After 7–10 days, the swelling gradually subsides. There may be an associated acute orchitis, which may be bilateral and which may proceed to testicular atrophy. A rare complication is pancreatitis.

**ACUTE SUPPURATIVE PAROTITIS**

Acute parotitis as a complication of major surgery is now quite unusual (Fig. S.1); this is because it results from postoperative dehydration in a patient with poor oral hygiene and septic dental stumps.

Nowadays, this condition is usually obviated by adequate fluid replacement and by both preoperative and postoperative oral care. Acute parotitis is also occasionally seen as a complication of the severe dehydration in conditions such as typhoid fever and cholera. Radiotherapy to the parotid region may result in damage to the gland, reduction in its secretion and a propensity, therefore, for ascending infection to occur.

On examination, the whole of the gland is enlarged, with a tender red swelling of the side of the face. This may progress to overlying cellulitis. Pus can be expressed from the parotid duct on the affected side. Because of the dense overlying fascia, which confines the enlarged gland, pain may be intense. There are the associated features of severe infection with pyrexia and toxaemia.

Figure S.1 Acute postoperative right-sided parotitis following gastrectomy for gastric carcinoma.
PAROTITIS ASSOCIATED WITH DUCT OBSTRUCTION

Obstruction of a salivary duct, from any cause, results in a typical syndrome in which the gland becomes painful and swollen at meal times, due to the increased secretion of saliva being unable to discharge through the duct. Between meals, as the saliva gradually escapes, the swelling and pain subside. Frequently, inflammation of the obstructed gland occurs as a result of ascending infection from mouth organisms. In these circumstances, there may be the associated features of infection (Fig. S.2) and there may be a discharge of pus from the mouth of the duct.

Although less common than in the submandibular duct, calculi in the parotid duct are not rare. They tend to be smaller and less radio-opaque than in the submandibular duct or gland, so that only larger ones are seen on a plain X-ray of the region. A sialogram may be necessary to identify the stone, which will then be seen as a filling defect.

Other causes of parotid duct stenosis are trauma from the irritation of an adjacent tooth stump or, occasionally, from traumatic division of the duct, for example following a knife laceration of the cheek.

SUBMANDIBULAR CALCULUS

Calculi are the most common cause of a painful swelling of the submandibular gland, and these account for some 95 per cent of all salivary stones. There are several reasons for the comparative frequency of stones in the submandibular gland and its duct. Its secretion contains more mucus than the parotid duct, so that it is more viscid. Its duct is longer and slopes upwards from the gland, so there is more tendency for a small concretion to remain within the duct. Furthermore, its orifice, being on the floor of the mouth, is more exposed to trauma than that of the parotid duct. The aetiology of these stones remains the subject of debate. Their size varies from minute to the size and shape of a date stone. Numbers vary: there may be a single stone in the duct, or the gland itself may contain numerous stones throughout a dilated duct system (sialectasia).

The classic story of a swelling and pain associated with food is elicited in the history. The gland itself can be felt as a tender enlargement on bimanual palpation with one finger below the angle of the jaw and the other in the sulcus between the tongue and the mandible. Occasionally, a calculus can be seen to extrude through the duct orifice at the side of the base of the fraenum linguae. At other times, it may be palpated along the course of the duct in the floor of the mouth, or stones may be felt in the gland itself. Pus may be expressed from the duct orifice by pressure on the gland.

Submandibular calculi are nearly all radio-opaque and can be visualized on a plain X-ray of the floor of the mouth. If there is difficulty in visualizing the calculus, it can be demonstrated as a filling defect on a sialogram of the duct (Fig. S.3).

CARCINOMA OF THE SALIVARY GLANDS

In its late stages, a carcinoma of the salivary gland (most common in the parotid, less often seen in the submandibular gland, and rare in the sublingual gland and the accessory salivary glands) will invade adjacent tissues and produce severe pain. Invasion of a salivary gland from an adjacent tumour, for example a carcinoma of the floor of the mouth, a squamous carcinoma of the overlying skin or a malignant melanoma, will be associated with intense pain.
The salivary glands are subject to swelling due to inflammation and new growth in the same way as any other organ. In common with other externally secreting glands, they are also subject to swelling resulting from retention of secretion. This most commonly occurs as a result of blockage of a duct by a stone. Parotid swelling with fever, often with lacrimal adenitis and uveitis (Mikulicz's syndrome), may occur in leukaemia, Hodgkin's disease, tuberculosis, systemic lupus erythematosus and sarcoidosis. Confusion in diagnosis may result from the close proximity of the lymph nodes; in the case of the submandibular gland, the lymph nodes may be right in the centre of the salivary tissue. The different salivary glands do not exhibit the same liability to each lesion, the submandibular for instance being the most liable to calculus formation, while inflammatory lesions are only common in the parotid. Mumps is the most common cause of all parotid swellings; it may occasionally involve glands other than the parotid, but this is a rare exception and usually occurs only after the parotid is first attacked. Here, as in all diagnosis, it is important to decide the exact anatomical site of the lesion before considering its pathology. For example, swelling of the loose tissues over the jaw from alveolar inflammation may mimic parotitis. A useful point in this connection is that a generalized parotid swelling tends to lift the auricle away from the head, and inspection of the orifice of Stensen's duct within the mouth will usually reveal some abnormality. If lymph nodes are suspected as a site of swellings, the presence of other enlarged nodes and of a primary lesion should be sought. Sialography may prove helpful. Radio-opaque contrast is injected into the appropriate orifice (Wharton's submandibular or Stensen's parotid; the lingual ducts are not suitable for injection). The branching system of ducts is well visualized in the radiograph. Blockage by a stone or by growth, sialectasia (dilatation and beading of the duct system, especially in the parotid gland) or the presence of a fistula is the lesion most likely to be demonstrated in this way. The lesions of the salivary glands are summarized in Table S.1.
Salivary tumours can be classified as:

- **Benign**
  - Pleomorphic adenoma (‘mixed tumour’)
  - Adenolymphoma (Warthin’s tumour)

- **Malignant**
  - Primary – carcinoma
  - Secondary
  - Invasion from overlying skin (e.g. malignant melanoma)
  - Secondarily involved lymph nodes

Approximately 90 per cent of **pleomorphic adenomas** (Fig. S.4) occur in the parotid, although the lesion is occasionally found in the submandibular gland (Fig. S.5) and, rarely, in the sublingual and accessory salivary glands. Likewise, 90 per cent are present before the age of 50 years, and the gender distribution is equal.
Characteristically, the tumour arises as a lobulated firm mass, noticed first when about the size of a cherry, and of variable consistency. The lump is painless and is typically situated between the ascending ramus of the mandible and the mastoid process, although no part of the parotid is exempt from this change, and these tumours may be found as low as 2.5 cm below the angle of the mandible. A frequent history is that the lump, over a period of years, shows a progressive slow increase in size.

Adenolymphoma accounts for about 10 per cent of parotid tumours and is very rare elsewhere. It usually occurs in men over the age of 50 years, and it may be bilateral. The tumour feels soft and cystic. Carcinoma (Fig. S.6) again usually affects the parotid, but it may occur rarely in the other main, and accessory, salivary glands. Usually, the patient is over the age of 50 years. Clinically, the diagnosis is based on rapid growth, pain, and involvement of the facial nerve (in the case of the parotid) and the regional nodes. Eventually, the surrounding tissues are infiltrated, and the overlying skin becomes ulcerated.

SARCOIDOSIS

In sarcoidosis, asymptomatic enlargement of the parotid, sublingual and submandibular glands occurs in about 6 per cent of cases. Spontaneous resolution often occurs. The glands are not tender. Facial palsy may occur with parotid enlargement. The syndrome of fever, uveitis and lacrimal and salivary gland enlargement is known as ‘uveo-parotid’ fever.

TINEA CAPITIS

Scalp ringworm (tinea capitis) is a fungal infection of the scalp, and predominantly a disease of childhood. It is contracted by direct contact. It had almost disappeared in the United Kingdom by the mid-1980s but has once again reached almost epidemic proportions, predominantly in subjects of Afro-Caribbean origin. The infection may be caused by a number of species of fungus; some such as *Trichophyton violaceum* and *Microsporum audouinii* are anthropophilic, and may thus be spread from child to child, including through fomites such as school caps, hair brushes, or barbers’ clippers. Other species, such as *Microsporum canis*, are zoophilic and are usually acquired by contact with animals, especially household pets. *Trichophyton verrucosum*, the causative organism of cattle ringworm, may cause a particularly violent inflammatory infection in children, and should be suspected if there is a history of contact with cows. In some cases of scalp ringworm, the inflammatory reaction is so marked as to produce a boggy discharging mass, termed a ‘kerion’, and is usually associated with local lymphadenopathy; if not promptly recognized and treated, this will progress to scarring and permanent hair loss, and such cases are a common cause of litigation.

In cases of scalp ringworm, the hair follicles are first infected, causing small, red, scaly patches in which hairs break off short, rather like down-trodden stubble (Fig. S.7). Lesions may remain solitary or be multiple.
Itching may or may not be present, and the degree of scaliness varies from a fine, branny desquamation to heaped-up masses of soft scale. Diagnosis is confirmed by microscopic examination of a few stumps that have been soaked in 10 per cent potassium hydroxide, but specimens should also be sent for laboratory culture, because identifying the species of fungus is important in determining the likely source. Scalp examination using an ultraviolet lamp (Wood’s light) can be most helpful, especially in screening groups of patients, as infections with small-spored fungi cause green fluorescence (Table S.2).

Tinea capitis must be differentiated from seborrhoeic dermatitis, where there may be diffuse fine scaling but no broken-off hairs. In scalp psoriasis, there are well-demarcated areas of scale heaped over red plaques through which the hairs grow undisturbed. In alopecia areata, there are one or more sharply demarcated bald areas entirely devoid of scaling or inflammation.

TINEA FACIEI
Ringworm on the face is likewise usually seen in children as the result of exposure to an animal harbouring a zoophilic species of fungus.

TINEA BARBÆ
Ringworm of the beard area is now rarely seen. It is largely a disease of the adult male and confined almost exclusively to agricultural workers who contract the disease from infected animals. It may take the form of superficial patches with folliculitis, or more usually a deep suppurative type (kerion) (Fig. S.8). Trichophyton verrucosum (cattle) and Trichophyton mentagrophytes (hedgehogs) are responsible for the majority of cases. Anthropophilic species are occasionally causes, and Microsporum canis can affect eyebrows and eyelashes.

### Table S.2 Fungus infections of scalp or beard

<table>
<thead>
<tr>
<th>Fluorescence with Wood’s light</th>
<th>Clinical features</th>
<th>Usual geographical location</th>
</tr>
</thead>
<tbody>
<tr>
<td>Small-spored</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Microsporum audouinii</td>
<td>+</td>
<td>Scaling bald patches</td>
</tr>
<tr>
<td>M. canis</td>
<td>+</td>
<td>Inflammatory scalp ringworm</td>
</tr>
<tr>
<td>Taenia violaceum</td>
<td>–</td>
<td>Black dot fungi</td>
</tr>
<tr>
<td>T. schoenleinii</td>
<td>–</td>
<td>Favus with scutuli</td>
</tr>
<tr>
<td>Large-spored</td>
<td>–</td>
<td>Scaling bald patches</td>
</tr>
<tr>
<td>T. tonsurans</td>
<td></td>
<td>Occasionally kerion</td>
</tr>
</tbody>
</table>

Figure S.8 Kerion (inflammatory ringworm).

Barry Monk

Scaly eruptions are thin plates of keratinized epithelial cells that occur as a result of an alteration of the normal keratinization process. Normally, as the epithelial cells become slowly impregnated with keratin, their nuclei are lost; eventually, they desquamate invisibly from the surface and are responsible for the majority of house dust. When the process is abnormally speeded up, keratinized cells may reach the surface of the skin before losing their nuclei, and adhere together in abnormal clumps; these appear white or silvery rather than translucent, and are shed as visible scales. This can occur in many chronic inflammatory conditions of the epidermis, particularly psoriasis. When the rate of epithelial turnover is markedly slowed (e.g. in myxoedema, severe malnutrition or ichthyosis), keratinized cells may accumulate abnormally at the skin surface, which again loses its translucency and is shed in large abnormal pieces.

Ichthyosis is an inherited disorder in which there is a generalized scaling of the skin without associated inflammation. Acquired ichthyosis should raise the suspicion of an underlying malignancy.

The major causes of scaly skin conditions are listed in Box S.1.

### RED SCALY LESIONS

Psoriasis is a common disorder of keratinization, which may start at any age, and is believed to have a genetic basis; a family history can be elicited in about 50 per cent of cases but without any consistent pattern of inheritance. It presents with sharply demarcated
thickened red plaques, surmounted by an easily detachable silvery scale (Fig. S.9). When affected areas are rubbed, punctate haemorrhage may be noted on the affected surface. In flexural areas, scaling may be absent, but there is a sharply demarcated glazed erythema. All areas of the body – from the top of the scalp to the soles of the feet – may be affected, but there is a predisposition to extensor surfaces and sites of friction. Lesions may localize in sites of injury to the skin, such as cat scratches or surgical wounds (the Koebner phenomenon). The scalp is commonly affected, but hair loss is uncommon. The extent of the eruption may vary from the inconsequential to whole-body involvement. When very extensive, systemic disturbance may arise through excessive vasodilatation of the skin. The nails are involved in about 50 per cent of cases, and there may be an associated arthropathy. Psoriatic plaques may go into spontaneous remission and, when they do so, they commonly clear from the centre, leaving an annular pattern.

In psoriasis, the lesions are generally multiple, and the presence of a solitary lesion of psoriasis should cause one to question the diagnosis. There may in fact be lesions in occult sites such as the scalp or ears; if truly solitary, the possibility of Bowen's disease (Fig. S.10) or a superficial basal cell carcinoma should be considered. Keratoderma blenorrhagica is a cutaneous reaction pattern comprising thickly scaled, red plaques, often in bizarre shapes and occurring chiefly on the soles, palms and genitalia. The lesions have a great similarity to psoriasis, but they follow urethritis and are accompanied by uveitis, fever, arthralgia and sacroiliitis. This symptom complex is called reactive arthritis, and many authorities believe that it occurs only in those with the psoriatic genotype. Another distinctive variant is guttate psoriasis, in which a streptococcal infection is followed by the sudden onset of a profuse eruption of tiny psoriatic lesions. The condition must be differentiated from pityriasis lichenoides chronica, which affects an older age group, does not follow a streptococcal sore throat, and where individual papules are covered by a single translucent ‘mica’ scale that detaches en masse. The lesions are also less numerous, and in varying stages of evolution at different sites.

Chronic eczema may also manifest itself with multiple red, scaly plaques. In discoid (nummular) eczema, there are multiple, intensely itchy, coin-shaped lesions scattered randomly over the trunk and limbs. This form of eczema is most commonly seen in middle-aged men and is commonly provoked by stress. Eczematous lesions are markedly less sharply demarcated than those in psoriasis, and the scales are usually finer and more adherent.

Eczema is not, of course, a diagnosis, but rather a pattern of reaction that may arise from a number of causes such as contact allergy, chronic exposure to irritants, photosensitivity or endogenous factors.
The pattern of the eruption may give a clue to the cause; for example, a localized area of eczema under a wristwatch may be caused by nickel sensitivity. In any case of unexplained eczema, investigation by patch testing should be undertaken if possible.

**Seborrhoeic dermatitis** is a distinctive pattern of endogenous eczema in which rather greasy superficial scales arise on a background of erythema; typically, the eruption is symmetrical, and affects the scalp, eyebrows, nasolabial folds and presternum. **Intertrigo** is a term used to describe any flexural rash, especially of the submammary folds and groins; many cases are due to seborrhoeic dermatitis, sometimes with superimposed candidiasis.

As with psoriasis, the possibility of a solitary scaly patch of eczema should be regarded with some suspicion, as this may in fact be a presentation of Bowen's disease or superficial basal cell carcinoma. However, one exception to this rule is provided by **lichen simplex chronicus**; this is a form of neurodermatitis caused by habitual scratching, and it presents with a thickened scaly plaque in which the skin markings are accentuated. The ankle is a common site in males, and the nape of the neck in women.

In **pityriasis rosea**, an acute eruption of unknown cause, the onset is often marked by the development of a solitary scaly oval lesion on the trunk (the herald patch), followed some days later by a widespread eruption of scaly ovoid lesions over the bathing trunk area. The degree of pruritus is very variable, and the rash resolves in 6–8 weeks. Inspection of individual oval lesions reveals the scale to be concentrated in a collarette just inside the edge. If lesions extend to the palms and soles and are accompanied by malaise, lymphadenopathy, fever and a sore throat, secondary syphilis should be suspected. The plaques are less pink and more of a ham colour. There is often patchy alopecia and ‘snail-track’ ulceration of the oral mucosa, the serology will be positive, and a primary chancre may still be present.

**Pityriasis versicolor** is a common superficial yeast infection that is generally seen in young adults (Fig. S.11). The individual lesions are symmetrically arranged over the neck and upper trunk, and are composed of flat, oval macules with a slight scale and minimal inflammatory reaction. Microscopy of the skin scrapings will reveal the causative organisms.

**Dermatophyte** infection can cause isolated red scaly plaques, usually with a central clearing and a raised red edge with an advancing scaly border; zoophilic fungi show considerable inflammation and rapidly expanding rings, anthropophilic species being more indolent (Fig. S.12).

**Mycosis fungoides** (cutaneous T-cell lymphoma) generally presents with a slowly progressing rash that may in some places resemble eczema and in others psoriasis. It is distinguished from both by its characteristically bizarre arcuate patterning, sometimes with islands of sparing. There may be rather variable erythema and scaling with a notable lack of response to topical therapy. Over the course of many years, the lesions extend, become more indurated, and may ultimately progress to tumours and systemic dissemination. Occasionally, cases present with tumours from the outset (**tumour d’emblée**).

**SCALY PALMS**

A unilateral scaly palm is the hallmark of **tinea manuum**. The colour is of a dull, lustreless greyish-red, and the scaling is fine and concentrated in skin creases. There may be associated fungal onychodystrophy of one or many fingernails and signs of tinea elsewhere.
on the body. The condition is often of very long standing. Bilateral scaly symmetrical palms can be inherited as a familial trait in the extremely rare tylosis palmaris. The hyperkeratosis is tightly packed and yellowish in colour. Similar yellowish, thick hyperkeratosis of palms is seen in pityriasis rubra pilaris, an uncommon condition that may be seen at all ages and may be accompanied by a widespread psoriasiform rash with follicular accentuation.

**SCALY SCALP**

The scalp is one of the sites of predilection of psoriasis, where well-demarcated plaques of thick scale cause surprisingly little disturbance of hair growth. The shedding of large amounts of silvery scales from these lesions is often a source of great embarrassment for patients. Seborrhoeic dermatitis causes a more confluent scaly eruption, and the scales are greasier and more adherent. Corroborative evidence should be sought of seborrhoeic dermatitis in the classic areas elsewhere on the skin. Tinea capitis usually causes asymmetrical scaling eruptions of the scalp. The hairs are usually broken off close to the surface, giving a stubbled appearance.

**SCALING OF WHOLE SKIN SURFACE**

Inflammatory skin conditions can, on occasion, affect the whole skin surface causing an erythroderma; when large amounts of scale are shed, the term exfoliative erythroderma is applied. This may develop from extension of a previously diagnosed skin condition (e.g. psoriasis), or present in this manner, when the diagnosis may be more difficult to establish. Possible causes include drug allergy (e.g. gold, antimalarials or sulphonamides), severe contact dermatitis, atopic dermatitis, seborrhoeic dermatitis, psoriasis, underlying lymphoma and Sézary syndrome (the erythrodermic stage of mycosis fungoides).

Erythroderma is often associated with significant constitutional upset, with difficulty of thermoregulation and a loss of fluid and protein. It has a significant mortality.

**SCROTUM, SURFACE AFFECTIONS OF**

Barry Monk

The skin of the scrotum is structurally different in that it is very thin, highly innervated and with a profuse blood supply (to allow local cooling). These characteristics make it prone to certain dermatoses, and any skin disorder affecting the area may be intensely itchy. The scrotum may be affected as part of a generalized eruption, such as seborrhoeic dermatitis or psoriasis. It may be the only area affected, and skin disorders in this site may be influenced by the local environment – heat, humidity and mobility of the part, local friction, and the thinness of the scrotal skin. In a bed-ridden patient, it may be irritated by urine or faeces, or the drugs they may contain (e.g. danthron breakdown products); an analogous situation may arise in infants allowed to stay in soiled nappies (ammoniacal dermatitis). Acute dermatitis and intertrigo are common on scrotal skin (Box S.2).

Acute dermatitis of the scrotum frequently follows over-treatment; the thin skin is particularly sensitive to medicaments such as those used in the treatment of pediculosis pubis or tinea cruris. Topical steroids penetrate the scrotal skin more readily than other sites, and this may lead to atrophy, and increased sensitivity of the scrotal skin. Seborrhoeic dermatitis may affect the scrotum and be confused with candidiasis (Fig. S.13) or with tinea. The latter nearly always affects the neighbouring groin skin, has a well-defined wavy edge, and shows mycelia when the scales are examined under the microscope. Moreover, patches of
white macerated or scaly skin produced by the same fungus can often be found elsewhere (e.g. between the toes). It is when a seborrhoeic dermatitis of the scrotum and groin is mistakenly treated for tinea (e.g. with Whitfield’s ointment) that a particularly acute and painful dermatitis can occur.

*Erythrasma* occurs as a discoid plaque of a beefy, brownish-red colour covered with fine scales; unlike tinea, there is no central healing and the plaques are uniform in appearance. They fluoresce pink under Wood’s light, and culture reveals the causative *Corynebacterium minutissimi*. Plaques of *psoriasis* can affect the scrotum, especially in flexural psoriasis. Scaling is less prominent at this site, but plaques will have the typical well-defined edge and salmon-pink colour. There should be evidence of psoriasis elsewhere on the body (e.g. the elbows, knees, sacrum and scalp).

*Syphilis* gives rise to moist papules or serpiginous, erythematous patches and ulcers; however, in contrast with dermatitis, the whole of the scrotal skin is not affected, itching is absent, and the serology will be positive. Hard and soft chancres may affect the scrotum as well as other parts of the genitals; the differential diagnosis is discussed under SCROTUM, ULCERATION OF (p. 604).

Intensely itchy nodules on the scrotum should suggest the diagnosis of scabies, and the diagnosis may be confirmed by finding the characteristic burrows on the wrists, fingers and elsewhere. Sometimes the scrotal nodules may persist as a post-scabetic phenomenon when the rest of the eruption has resolved with treatment.

Boils, warts, sebaceous cysts and tumours are referred to under TESTICULAR SWELLING (p. 666). Some scrotal neoplasms may cause non-healing erosive and ulcerating lesions; examples include *Bowen’s disease*, *basal cell carcinoma* and rarely *extramammary Paget’s disease* (Fig. S.14). Moles and malignant melanomas can also occur on scrotal skin.

Ulceration of the scrotum may be a feature of *Behçet’s syndrome*, the features of which are recurrent orogenital ulceration, together with iritis, keratitis, diffuse pustulation, polyarthritis, thrombophlebitis and sometimes neural infarcts.

Small angiomas are very common on the scrotum in elderly persons and are known as *angiokeratomas of Fordyce*. Small nodules on the scrotum and penis are characteristic of *scabies*. Discharging nodular lesions on the scrotum, spreading into the groins and often affecting the axillae, are seen in *hidradenitis suppurativa.*
Recurrent erosions in the same site can be caused by both herpes simplex and fixed drug eruptions. In the former case, multiple vesicles and minimal scarring will be seen. In the latter case, considerable post-inflammatory hyperpigmentation is the rule, and a carefully taken drug history will usually reveal the culprit (e.g. codeine, laxatives or sulphonamides).

In lichen simplex chronicus, the habit of regular rubbing and scratching leads to considerable thickening and lichenification of the skin; this is common in perianal skin, but it also occurs on the scrotum, usually in an intense and nervous subject; treatment may be challenging.

### SCROTUM, ULCERATION OF

**Ben Challacombe**

Ulceration of the scrotum occurs in association with:

- Underlying disease of testis
  - Inflammatory (epididymo-orchitis)
  - Tuberculous
  - Syphilitic
  - Neoplasm
- Fistula
  - Syphilis involving the scrotum
  - Suppurating cutaneous cysts
  - Infected haematocoele
  - Cutaneous neoplasms
- Behçet’s syndrome, herpes simplex and candidiasis

### UNDERLYING DISEASE OF THE TESTIS

In some cases, extension of disease in the testicle may involve the coverings of the scrotum, and it may even perforate them to form a scrotal sore. This sequence occasionally occurs with:

- Testicular abscess
- Severe Epididymo-orchitis
- Tuberculosis of the epididymis
- Gumma of the testis
- Malignant disease of the testis

A testicular abscess is somewhat uncommon but may arise from direct extension from the urethra via the seminal vesicles and vasa deferentia, or by a haematogenous infection during the course of a specific fever, such as scarlet fever, mumps or typhoid fever. With urethral disease, the primary problem may be due to gonorrhoea, or more frequently to a septic urethritis from the introduction of infected instruments. In cases in which the infective process extends from the urethra, the epididymis is affected first, whereas in the metastatic cases, the body of the testis usually shows the first sign of enlargement.

On occasion, following vasectomy for sterilization, the swelling and possible abscess will occur in the upper part of the scrotum at the site of the division. Pathology causing acute inflammation of the testis occasionally suppurates, when the scrotal tunics become inflamed and adherent, while softening occurs later. Unless surgically relieved, the abscess opens through the skin, leaving an ulcer and a sinus discharging pus. An unusual form of abscess of the testicle is caused by a suppurating dermoid cyst of the testicle, and this may discharge through the scrotal coverings to form an ulcer.

Tuberculosis of the testis rarely occurs as a primary disease but more often as a secondary deposit in association with tuberculosis elsewhere in the genitourinary tract. Testicular tubercle almost always begins as a nodule in the epididymis, but in the later progress of the disease may extend into the testis proper. If the tuberculous nodule progresses rather than undergoes cure, the scrotal skin becomes adherent, thinned and finally perforated, leaving a shallow ulcer with thin, undermined edges, and discharging thin pus. The ulcer in this case is most likely to be on the posterior aspect of the scrotum. Occasionally, the necrotic epididymis fungates through the opening in the scrotum, appearing as a greyish, sloughy projection from the cutaneous opening – the so-called ‘hernia testis’.

A gumma of the testis (once common, now very rare), causes a swelling in the body of the testis rather than in the epididymis. A gumma that remains unrecognized or untreated may soften and ulcerate through the scrotal skin in a manner similar to tuberculous disease, leaving a clearly defined ulcerated area with sharply cut margins and a wash-leather-like, sloughy base. Such ulcers are usually placed on the front of the scrotum. The gummatous granulation tissue may fungate through the scrotal aperture, forming a yellowish necrotic mass.

The diagnosis of these three conditions may produce some difficulty in the earlier stages (see TESTICULAR SWELLING, p. 666), but, in the advanced stage now under consideration, when an open scrotal sore is present, the diagnosis is easier.

The opening of a testicular abscess on the scrotum leaves a small sinus discharging pus and accompanied by a general enlargement of the organ. Preceding the rupture of the abscess, there is acute pain in the testicle, with a rise in temperature, rigors and general signs of suppuration, although these are
much diminished as soon as the abscess bursts or is incised. There is often a urethral discharge, but this is frequently much lessened with the onset of acute epididymitis, with distinct thickening of the cord and aching pain in the region of the external abdominal ring. In metastatic cases, the abscess occurs during the progress of an acute fever. The general history is one of acute pain beginning in the testicle, with rapid and extremely tender swelling of the organ, followed by abscess formation.

In tuberculosis of the epididymis, the progress is much more gradual. A nodule may have been present in the epididymis for some time, gradually enlarging, but causing very little pain. In some cases, a nodule may have been present for months without any apparent change, and then it may enlarge rapidly, involve the scrotal tunics, and discharge its contents. By the time the disease has reached this stage, it is probable that evidence of tuberculosis will be found in other organs, particularly the other testicle, prostate, seminal vesicles or bladder. The affected testicle usually presents several nodules in the epididymis and is tender on pressure, while small nodules may also be felt in the vas deferens.

The opening remaining from the discharge of a gummatous orchitis is usually a rounded ulcer with sharply cut edges and a yellowish base. The whole testis is enlarged and practically painless. The cord is not thickened, and there is no evidence of disease in the other testicle, prostate or seminal vesicles. There is probably a history of syphilis, and other tertiary syphilitic lesions, such as gummatous periostitis, may be present elsewhere.

Neoplasms of the testis seldom cause ulceration of the scrotum because they have generally been removed surgically before such a late stage is reached. Any variety, however — whether seminoma or teratoma — may cause local recurrence in the scar, with ulceration. The diagnosis depends upon histological examination, either of the tumour previously removed or of a biopsy from the edge of the recurrence. Occasionally, the fungation of the tumour is seen through the scar of the biopsy site in the scrotal skin when there has been delay in carrying out definitive treatment, although this state of affairs should never be allowed to happen.

**FISTULA**

Fistulas may occur in the scrotum and cause ulceration. Sinuses occur in association with tuberculosis or syphilitic disease of the testes, but fistulas may follow urine extravasation, or from rectal suppuration. An abscess may form and open through the scrotal skin from a peri-urethral abscess accompanying an acute urethritis or formed by septic infection behind a urethral stricture. In either case, a small amount of urine may leak through the opening during micturition, while the history of the urethral discharge, or of difficulty in micturition and other symptoms of stricture, will point to the diagnosis.

**SYPHILIS INVOLVING THE SCROTUM**

This may be present either as a primary chancre or as a mucous tubercle. A primary chancre in this situation is by no means easy to recognize unless other signs of syphilis are present. However, the presence of a cutaneous sore that does not show much inclination to heal under antiseptic dressings should always give a suspicion of syphilis. There is often only slight induration of the ulcer compared with that of a penile chancre, but the edge is raised and of a rolled appearance. The inguinal lymph nodes are enlarged and discrete, but it is some 5–6 weeks after the commencement of the ulcer that the usual secondary symptoms of syphilis become manifest.

Mucous tubercles of secondary syphilis may be present on the scrotum, usually on the lateral aspect. They may extend directly from the anal area. No difficulty will be met with in the diagnosis, as other signs of syphilis are obvious and the specific serological tests are positive.

**SUPPURATING CUTANEOUS CYSTS**

A sebaceous cyst in the scrotal skin may suppurate and leave an open sore. The areas remaining present raised borders, and are easily mistaken for an early squamous cell carcinoma. An accurate history of the previous swelling in the skin is of little assistance in these cases, but microscopic examination of a piece removed from the margin of the ulcer will exclude malignancy.

**CUTANEOUS NEOPLASMS**

Carcinoma of the scrotum, formerly known as ‘chimney-sweep’s cancer’ or ‘tar-worker’s cancer’, is by no means limited to these occupations, but related to chronic irritation from solid particles or from noxious fumes. Hence the disease is, or was, most commonly seen among chimney sweeps, employees in gas works, paraffin, tar and chemical works and coal mines, and in spinners in the cotton trade. It often begins as a small subcutaneous nodule, over which the skin is thinned and adherent. The nodule enlarges slowly, and the thinned covering gives way to form an ulcer with thickened irregular edges and a tendency to bleed on slight injury. The ulcerated area extends both radially and into the tissues of the scrotum, later involving the testes. The inguinal nodes become enlarged soon after active ulceration begins, at first from inflammatory...
causes, but later from malignant infiltration. If left untreated, these nodes will ulcerate, and indeed the common mode of death in this condition is from repeated haemorrhages due to erosion into large vessels.

In other cases, a scrotal epithelioma begins in a wart or papilloma, which may have been present for years with only a slight increase in growth (Fig. S.15). These soft papillomas are not unusually the starting point of malignant change, when they become more vascular, while the surface epithelium becomes thinned and easily excoriated. A small amount of foul discharge is present, often encrusted into a scab that, on removal, leaves an ulcer with indurated, everted edges, with the gradual progress of a cutaneous squamous cell carcinoma.

Any ulcer on the scrotum – especially if indurated or readily caused to bleed – must be looked upon with extreme suspicion and immediately subjected to biopsy for microscopic examination. It is not unusual, however, for a large mass of nodes to be found in the groin when the primary lesion is very small and almost imperceptible. The scrotum must be examined very carefully in such cases lest the primary lesion be missed.

**Figure S.15** Epithelioma of the scrotum.

**BEHÇET’S SYNDROME, HERPES SIMPLEX AND CANDIDIASIS**

Behçet’s syndrome causes painful ulcerative lesions of the scrotum as well as the penis, unlike the lesions in the vulva and vagina, which are often painless and therefore often missed. Behçet’s may be accompanied by abscess or herpes-like lesions of the scrotum. Herpes simplex – both types I and II – may cause vesicular lesions less commonly, and very rarely candidiasis.

**SELF-HARM, DELIBERATE**

Andrew Hodgkiss

Self-harm remains the most common reason for emergency hospital care in young adults, accounting for 10 per cent of all acute admissions, or about 170 000 cases each year in England and Wales. The term encompasses non-accidental self-poisoning and non-accidental self-injury (such as cutting, jumping, shooting or hanging). ‘Self-harm’ is a term reserved for those who survive such acts, *completed suicide* being the word used for fatalities.

The initial management is of course the medical and/or surgical evaluation and treatment of the physical presentation. Psychosocial assessment of the behaviour takes place subsequently when the patient is fit to be interviewed. The first psychological task is to confirm whether non-accidental self-harm actually took place, and to accomplish this, the account of an individual close to the patient is always desirable and may prove essential. The types of self-harm are listed in Box S.3.

*Feigned* self-poisoning is by no means rare, suspicions being raised by atypical symptoms, responses or behaviour and the absence of confirmatory physical signs or evidence. Such patients may be habitual self-poisoners who eventually only go through the motions, or may be a self-referral from out of town with a dramatic story and no corroborative evidence available, when the possibility of Munchausen’s syndrome should be considered.

**Box S.3 Types of self-harm**

**Most common**
- Deliberate

**Less common**
- Experimental
- Accidental
- Feigned
- Self-mutilation in the learning disabled
Experimental self-poisoning is usually evident from the drugs taken and the account of somebody present at the time. Most common are illicit substances such as opiates, barbiturates, amphetamines, magic mushrooms and solvents, but prescribed drugs, particularly minor tranquillizers, hypnotics, analgesics, anticholinergics and inhalers, are taken for this purpose, and proprietary medicines such as Actifed® and Benylin® may also be abused. It should not be uncritically assumed that the employment of an illicit substance necessarily implies that the purpose of the act was for recreation; the circumstances must also be taken into account.

Accidental self-poisoning can be harder to establish and, if doubt persists, the patient should be considered to have acted deliberately. This problem occurs most commonly when the patient’s judgement was impaired by alcohol, drugs or other physical causes of confusion, with an invariably sketchy recall of events. The necessity for an independent account is crucial in cases where accident is asserted by the patient, as this may be a ploy for evading recognition in an acutely suicidal person.

Self-mutilation in people with more profound degrees of learning disability is usually regarded as distinct in that these patients have diminished judgement, which raises doubts about their capability to form intent. However, self-injurious behaviour should never be assumed to be acceptable or normal in individuals with learning disability: its origins may lie in factors as diverse as epilepsy, bipolar disorder, under-stimulation and relationship problems within the family.

Having ascertained that non-accidental self-harm has occurred, the next step is to evaluate the act, and it is important to appreciate that self-harm is a common end point of behaviour, and not a medical or psychiatric diagnosis. The circumstances surrounding the act are a pointer to the degree of suicide intent, while the events leading up to taking the decision also help to establish the motive(s). It is crucial to understand that the majority of patients who deliberately harm themselves have no wish to kill themselves: other common motives include ridding oneself of unpleasant feelings, escaping from a stressful situation, obtaining help (the classic ‘cry for help’), and using it as a form of communication with another person (such as an expression of anger).

Next, precipitating and vulnerability factors should be inquired about in the patient’s social and personal history. Symptoms of mental illness, especially depression, should be specifically sought. Recent life events, especially losses and interpersonal difficulties, are particularly important. Factors that are associated with increased suicidal risk are presented in Box S.4.

Finally, mental state assessment should seek objective evidence of mental illness, as well as considering continuing suicidal risk. How the patient views their action and their future are essential inquiries; hopelessness is consistently reported to be strongly associated with subsequent self-harm and suicide.

This interview is often more difficult than in usual situations because it rarely takes place at the

<table>
<thead>
<tr>
<th>Box S.4 Profile of suicidal risk factors</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Circumstances surrounding the act</strong></td>
</tr>
<tr>
<td>• Evidence of planning and timing</td>
</tr>
<tr>
<td>• Evidence of precautions against discovery</td>
</tr>
<tr>
<td>• No communication or seeking help afterwards</td>
</tr>
<tr>
<td>• Suicide note, will, gifts, insurance changes</td>
</tr>
<tr>
<td><strong>Self-report about the act</strong></td>
</tr>
<tr>
<td>• Expectations of fatality</td>
</tr>
<tr>
<td>• Concepts of method’s lethality and reversibility (more important than quantity taken)</td>
</tr>
<tr>
<td>• Intention/motive(s)</td>
</tr>
<tr>
<td><strong>Sociodemographic variables</strong></td>
</tr>
<tr>
<td>• Male</td>
</tr>
<tr>
<td>• Middle-aged or elderly</td>
</tr>
<tr>
<td>• Widowed,* divorced,* separated* or single</td>
</tr>
<tr>
<td>• Unemployment*</td>
</tr>
<tr>
<td>• Social isolation or rejection*</td>
</tr>
<tr>
<td>• Financial difficulties*</td>
</tr>
<tr>
<td><strong>Physical variables</strong></td>
</tr>
<tr>
<td>• Any disabling, painful, chronic or life-threatening illness*</td>
</tr>
<tr>
<td>• Epilepsy</td>
</tr>
<tr>
<td><strong>Psychological variables</strong></td>
</tr>
<tr>
<td>• Family history of suicide</td>
</tr>
<tr>
<td>• Previous deliberate self-harm or psychiatric history</td>
</tr>
<tr>
<td>• Alcoholism or drug addiction</td>
</tr>
<tr>
<td>• Personality disorder</td>
</tr>
<tr>
<td>• Schizophrenia (especially during a quiescent phase)</td>
</tr>
<tr>
<td>• Depression especially if:</td>
</tr>
<tr>
<td>– Suicidal ideation/impulses</td>
</tr>
<tr>
<td>– Pessimism</td>
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<tr>
<td>– Self-denigration</td>
</tr>
<tr>
<td>– Hypochondriacal or guilt delusions</td>
</tr>
<tr>
<td>– Self-neglect</td>
</tr>
<tr>
<td>– Sleep disturbance</td>
</tr>
<tr>
<td>– Mood is cycling</td>
</tr>
<tr>
<td>– Early stages of treatment</td>
</tr>
<tr>
<td>• Hopelessness – may be the single most important predictor, irrespective of mood state</td>
</tr>
</tbody>
</table>

*Especially if this represents recent, abrupt change
patient’s request. Resentment at the suggestion of a psychiatric label, embarrassment about the behaviour with fear of the consequences, and unwillingness to disclose the true motive may all lead to poor compliance, while the lingering effects of alcohol or drugs may subtly impair the mental state. Nevertheless, it is important to persevere for three purposes: (i) the identification of treatable mental disorder; (ii) the identification of issues where intervention may help the patient and thus (hopefully) reduce the likelihood of recurrence; and (iii) the evaluation of suicidal risk.

Risk assessment requires an understanding of the factors associated with subsequent completed suicide, as well as the quite different list of risk factors associated with recurrent self-harm. While identifying those at high risk of completed suicide is medicolegally essential, it does not always lead to easy prevention of subsequent suicide, especially in those patients making life-threatening suicide attempts in the absence of severe mental illness. There is a growing research literature on the prevention of recurrent self-harm through psychological treatments that aim to improve patients’ problem-solving capability.

Sensation, abnormalities of

Mark Kinirons

Sensory disturbances are referred to in many sections of this book, but it is convenient here to consider the subject as a whole under the present heading.

Terminology

The inaccurate application of an exact terminology gives rise to so much confusion in the literature that it is often preferable to describe sensory experiences and sensory findings in plain language. ‘Paraesthesia’ is used for sensations of tingling, pins and needles, subjective numbness and feelings of cold and heat, whether they appear spontaneously or as a result of touching or manipulating the part. Since the term covers so many different sensations, it should be avoided when precise descriptive work is required, as for instance in case histories.

Anaesthesia means ‘without feeling’, but neurologists use it for loss of sensibility to light touch; partial reduction of such sensibility is called ‘hypoesthesia’ or ‘hypoesthesia’. Analgesia (loss of pain) and hypoalgesia or hypalgesia (reduction of pain sensibility) are useful terms, free from ambiguity. Thermanaesthesia and thermhypanesthesia (loss of, and reduction of, temperature sensibility) are explicit, if inelegant. These cutaneous sensibilities, together with those derived from the special senses, are called ‘exteroceptive’.

‘Proprioceptive’ sensibility is concerned with information received from the labyrinths and from muscle and joint receptors. Vibration sense is difficult to classify and is of no apparent value to humans (but it did warn our ancestors of the herd of animals they could not see but their naked feet could feel by vibration), but its loss may help to localize a lesion, especially of the spinal cord.

Hyperaesthesia, hyperalgesia and hyperthermaesthesia refer to increased sensibility to touch, pain and temperature, respectively. Increased sensibility to proprioceptive stimulation has not been described.

Anatomy and physiology

The four common cutaneous sensations – touch, pain, heat and cold – together with the deep sensations of pressure and proprioception – are referred to as the ‘somatic sensations’. These are consciously appreciated in all parts of the body, and they have a common pathway within the nervous system. An appropriate stimulus generates an impulse at the periphery that passes into the central nervous system, is relayed by the thalamus, and thence, by a final relay, is passed to the appropriate part of the cerebral cortex. In simple terms, the pathway for somatic sensation is subserved by three orders of neurones: the first-order neurone is concerned with transmitting information from the periphery to the spinal cord; the second-order neurone transmits information from the spinal cord to the thalamus; and the third-order neurone transmits information from the thalamus to the cerebral cortex.

Receptors

Information to the first-order neurones comes from a variety of receptors. Every sensation depends on impulses excited by the adequate stimulation of these receptors, which comprise two main groups – those in the skin, and those in the deeper somatic structures. Although individual cutaneous sensory receptors are most sensitive to a particular form of natural stimulation, this specificity is not absolute, and other forms of stimulation may also excite the receptors.

Transmission of information from receptors

The stimulation of a sensory ending gives rise to a receptor potential that appears at the specialized end of an afferent nerve fibre. This is not an all-or-none phenomenon but varies in amplitude and time course, and it may be rapidly dissipated even though the stimulus continues, with resulting falling away of the firing frequency in the nerve fibre. Impulses from the receptors travel centrally through the first-order neurones, the cell bodies of which are in the dorsal root ganglia.
Afferent pathways to the spinal cord

The afferent sensory fibres from the various receptors pass up the differing peripheral nerves and enter the spinal cord via the dorsal (posterior) roots. Understanding the variety of peripheral nerve disturbances that may occur requires knowledge of the anatomy of the distribution of peripheral sensory nerves and sensory roots. It should be emphasized that there is considerable overlap in the peripheral nerve distributions and in the dermatome distributions.

There is some evidence to suggest that a number of afferent fibres enter the cord in the ventral (anterior) roots. The significance of these is uncertain, but their presence may be responsible for the persistence of pain in some patients after dorsal rhizotomy.

Somatic sensory pathways

All of the somatic sensory pathways are crossed and terminate in the opposite sensory cortex in the cerebral hemisphere. Three anatomically separate pathways may be recognized: dorsal column medial lemniscus, lateral spinothalamic, and trigeminothalamic.

Dorsal column medial lemniscus pathway

Input to this pathway within the spinal cord is via large, thickly myelinated fibres that pass through the medial division of the dorsal spinal nerve root to enter the dorsal white column of their own side, dividing into ascending and descending branches. The descending branches establish reflex connections by sending collateral branches into the dorsal grey column; the ascending branches are the first link in the sensory pathway. At their entrance, these ascending fibres are situated immediately medial to the dorsal horn, but during their course up the spinal cord they are steadily pushed to a more medial position because the fibres entering at succeeding rostral (higher) levels intrude between the ascending fibres and the dorsal horn. As a consequence of this, the fibres occupying the most medial part of the dorsal column in the upper cervical region will belong to the sacral roots, while the fibres from the upper extremity are found most laterally. The fibres terminate at the cervicomедullary junction in the nucleus gracilis and nucleus cuneatus. The fibres terminate synaptically onto the second-order neurones in the gracile and cuneate nuclei, and the axons of these neurones curve ventrally and medially, crossing the midline, and then turn upwards to form a prominent bundle of fibres, the medial lemniscus.

The classical view is that the impulses ascending in the fibres of the dorsal columns mediate the sensations of touch, deep pressure, vibratory sense and sense of position of joints, and are particularly important for sensory discrimination. During the past few years, this view has been challenged and, although the matter is far from certain, it appears that the dorsal columns mediate sensory signals necessary for complex tasks.

The segmental somatotopic organization present in the dorsal columns and their nuclei is maintained in the medial lemniscus as it ascends to the thalamus, where the fibres enter the ventroposterior lateral nucleus, which contains the cell bodies of the third-order sensory neurones.

Lateral spinothalamic pathway

This pathway transmits impulses that are concerned with the appreciation of heat, cold and pain. It also provides an alternative pathway for touch sensibility – the so-called ‘crude’ or ‘coarse’ touch. The first-order neurones have their cell bodies in the dorsal root ganglia, and their fibres are thinner than those of the dorsal column medial lemniscus pathway; some, indeed (the C-fibres), have no myelin at all. They enter the spinal cord in the lateral part of the dorsal root, and divide into short descending and ascending branches. The ascending branches run for one or two segments in the posterolateral column before synapsing with second-order sensory neurones that lie deep in the dorsal column. The axons then cross the midline, the so-called ‘ventral white commissure’, and ascend in the ventrolateral white column as the spinothalamic tract. Some of the spinothalamic fibres give off collaterals to certain nuclear regions such as the reticular formation.

In the brainstem, the spinothalamic tract lies lateral to the medial lemniscus, which it accompanies to terminate in the thalamus in the ventroposterior lateral nucleus. Important features of the spinothalamic pathway include the following:

- The second-order neurone fibres cross the midline only one or two segments above the level of entry of the dorsal root fibres.
- The site of decussation (crossing) of the fibres in the cord exposes them to damage by expanding ventral cord lesions.
- Fibres concerned with pain and temperature sensibility are situated dorsally to those involved with touch and pressure.
- The spinothalamic tract is less compactly organized than the medial lemniscus, being intermingled with other ascending pathways giving off collaterals to the brainstem reticular formation.

Trigeminothalamic pathway

This pathway carries information from the distribution of the trigeminal nerve, which serves most of the skin of the face, the forehead as far as the vertex,
the mucous membranes of the nasal cavities, paranasal sinuses, mouth, tongue and parts of the pharynx, the teeth and gums, and part of the dura mater.

About half of the fibres entering in the trigeminal nerve divide into a branch that terminates in the chief nucleus of the trigeminal nerve, while the other half descends in the spinal tract to end in the spinal nucleus. The chief nucleus, which is in the lateral part of the pons, contains the second-order neurones concerned with tactile and postural sensibility; it gives rise to fibres that cross the midline to ascend near the medial lemniscus. The nucleus of the spinal tract, which extends downwards in the lateral part of the medulla to about the level of C2, contains the second-order neurones concerned with pain and temperature sensibility. Second-order neurones cross to the quintothalamic tract, which ascends close to the spinothalamic tract. Both sets of second-order neurone fibres terminate in the ventroposterior medial nucleus of the thalamus.

The thalamus and thalamocortical projections
The third neuronal link in the ascending somatic sensory fibre system is made up of neurones whose nuclei lie in the thalamus. The axons of these neurones transmit impulses to the cerebral cortex.

The somatosensory cortical areas
From clinical and physiological observations, it has been known for many years that, in man, the postcentral gyrus is the main (primary) somatosensory area. Another area beneath the lower end of the postcentral gyrus is known as the secondary somatosensory area. Since the work of Penfield, it has been known that there is a clear somatotopic representation in the sensory cortex. The first sensory area appears principally to reflect activity in the dorsal column medial lemniscus system and also in the associated trigeminal system. The thalamic relay for these impulses passes through the internal capsule.

However inadequate our understanding of the sensory cortex, one thing is clear: provided that the subcortical structures – especially the thalamus – are intact, certain sensations such as pain, touch, pressure and extremes of temperature can reach consciousness.

The accurate localization, however, as well as the patient's ability to make sensory discriminations, depends on the integrity of the sensory cortex. This is a fundamental distinction, and it will be discussed further when considering individual sensory syndromes.

PATTERNS OF SENSORY DISTURBANCE
Figures S.16–S.21 illustrate some of the sensory syndromes discussed below.
**Figure S.18** Dorsal myelitis affecting the cord as high as the ninth dorsal segment. The shaded parts are insensitive to touch, pain and all degrees of temperature.

**Figure S.19** Fracture–dislocation of the cervical spine. The shaded area represents the loss of sensibility to touch, pain, heat and cold.

**Figure S.20** Syringomyelia. The shaded parts show the areas of dissociated anaesthesia (i.e. of thermoanaesthesia and analgesia). This was associated with atrophic palsy of the upper extremities.

**Figure S.21** Thrombosis of the left posterior inferior cerebellar artery. The shaded areas show the regions of dissociated anaesthesia (i.e. loss of sensibility to pain and temperature of all degrees).
Mononeuropathy
Changes in this instance will vary, depending on whether the nerve involved is predominantly motor, sensory or mixed. In sensory nerves, the area of touch loss is usually more extensive than the area of pain loss. Because of overlap from adjacent nerves, the area of sensory loss following damage to a cutaneous nerve is always less than its anatomical distribution. Deep pressure and joint position senses remain intact because they are mediated by nerve fibres from the subcutaneous structures and joints. Particular types of pathological lesion may differentially affect the fibres in a sensory nerve. Compression typically disturbs large touch and pressure fibres, and leaves intact small pain, thermal and autonomic fibres. Lesions of the brachial or lumbosacral plexus may be differentiated from multiple peripheral nerve involvement by the distribution of the sensory and motor loss (see ‘Radiculopathy’, below).

Polyneuropathy
In most instances of polyneuropathy, the longest and largest fibres tend to be involved. The sensory loss is most severe over the feet and legs, and less severe over the hands; the trunk and face are usually spared except in the most severe cases. Typically, the sensory loss involves all the modalities, although this varies depending on the type of neuropathy. The term ‘glove and stocking’ sensory loss draws attention to the distal pattern of involvement (Fig. S.16). However, it is an inaccurate term as the border between normal and abnormal sensation is not sharp, and the sensory loss shades off gradually. In hysteria, the border between normal and abnormal sensation is usually sharp.

Radiculopathy
Irritative symptoms may be present when the dorsal roots are the subject of traction or compression. This shows itself as pain that is often limited to the dermatome belonging to the affected root. In some root disorders, pain is absent and paraesthesiae in the dermatome distribution are present. Damage to a dorsal root will result in a loss of sensory modalities of all types within the distribution of the dermatome. Because of overlap between the dermatomes, the interruption of one single dorsal root will often give no definite sensory loss. When two or more roots have been completely divided, the zone of sensory loss is usually greater for pain than for touch. Surrounding the area of complete loss will be a zone of partial loss.

SPINAL CORD SYNDROMES
Lesions of the dorsal horn (tabetic syndrome)
Lesions of the dorsal horn produce syndromes similar to those seen in lesions of the dorsal roots. Depending on the number of segments involved, there will be a segmental sensory loss affecting vibration and position senses in particular. Accompanying this may be pain, which often is called ‘lightning pain’. This repeated severe pain is described as occurring at right angles to the skin and penetrating through the affected limb. Most commonly, this syndrome results from neurosyphilis, although it may be seen in diabetes mellitus.

Transverse cord lesions
A complete transverse lesion of the spinal cord will be associated with a loss of all forms of sensation below the segmental level that corresponds to the lesion. There may be a narrow band of hyperaesthesia at the upper margin of the level of sensory loss. Loss of pain, temperature and touch sensation is usually evident two or three segments below the level of the lesion, whereas vibratory and position sense is less easy to delimit. A compressive cord lesion is usually associated with descending loss of sensation as the outermost fibres carrying pain and temperature sensation are from the legs. A lesion expanding from the centre of the cord, such as an intramedullary tumour, will tend to involve the innermost fibres carrying pain and temperature sensation, and thus there may be relative sparing of the most superficial fibres from the sacral segments; this may lead to so-called ‘sacral sparing’.

Hemisection of the spinal cord (Brown–Séquard syndrome)
Occasionally in spinal cord disorders, pathology is limited to one side of the spinal cord. Loss of pain and temperature sensation is found on the opposite side, and the upper margin of this is usually two or three segments below the level of the lesion. Proprioceptive sensation is affected on the same side as the lesion, and an associated motor paralysis occurs on the same side. Touch sensation is not involved, because the fibres are distributed in both posterior columns and the spinothalamic pathway on both sides of the cord. In clinical practice, a complete hemisection of the spinal cord is rarely seen, although a partial syndrome occurs in multiple sclerosis.

Central spinal cord lesions (syringomyelic syndrome)
A central spinal cord lesion will characteristically involve the pain and temperature fibres as they cross in the anterior commissure. Typically, these
modalities are affected on one or both sides over a number of dermatomes, with relative preservation of tactile sensation (so-called dissociated sensory loss). Abolition of tendon reflexes in the affected segment is usually seen. The most common cause of this syndrome is syringomyelia (see Fig. S.20), but intramedullary tumours (e.g. GHomas or ependymomas) may also produce it.

**Posterior column syndrome**

In lesions preferentially affecting the dorsal columns, there is loss of vibration and position sense below the level of the lesion, with preservation of pain, temperature and touch. When such sensory loss affects the legs, there is typically a sensory ataxia and a positive Romberg’s sign. Sensory loss of this type in the hands produces clumsiness when manipulating small objects, and an inability to recognize shapes such as coins in the pocket. Tingling and pins-and-needles sensations are common, and patients often complain that the hands and feet feel swollen or tight. Causes include vitamin B₁₂ deficiency and syphilis.

**Anterior cord syndrome**

In anterior cord disturbances, there is typically damage to the spinothalamic tracts, producing pain and temperature loss below the level of the lesion.

**Brainstem syndromes**

Because of the complex structure of the brainstem, with multiple ascending and descending tracts intermingled with a variety of cranial nerve nuclei, lesions result in far more complex clinical pictures than those which may be seen in spinal cord disorders. A characteristic feature of a medullary or lower pontine lesion is that the sensory disorder is crossed – that is, there is loss of pain and temperature sensation on one side of the face and on the opposite side of the body. This results from involvement of the trigeminal tract or nucleus, resulting in ipsilateral facial sensory loss, and of the lateral spinothalamic tract, resulting in a contralateral loss of sensation of the trunk and limbs. Higher in the brainstem, the trigeminothalamic and lateral spinothalamic tracts run together, and a lesion will therefore produce contralateral loss of pain and temperature sense on the whole of the opposite side of the body. In the upper brainstem, the spinothalamic tract and the medial lemniscus become confluent, so that a lesion at this level may cause contralateral sensory loss of all types.

In brainstem lesions, there is frequently bilateral involvement, and a variety of syndromes have been described involving sensory, motor and cerebellar dysfunction accompanied by cranial nerve palsies. Lesions, particularly vascular lesions, are rarely discrete, and it requires a detailed knowledge of the anatomical structure of the brainstem to achieve accurate localization. Another point of practical importance is that partial involvement of the sensory tracts may produce sensory impairment that may mimic lesions in the cord.

**THALAMIC DISORDERS**

Thalamic sensory disorders usually result from discrete cerebral infarcts. A destruction of the entire thalamic area receiving sensory fibre systems would be expected to result in an impairment or loss of somatic sensation in the whole of the opposite half of the body. In documented cases where such a lesion has been identified, the perception of pain has often been found to be only slightly affected. Position sense typically is affected more profoundly than any other sensory function. Pure lesions of the ventroposterior lateral nucleus of the thalamus will be associated with contralateral sensory disorders of the limbs and trunk, whereas involvement of the ventroposterior medial nucleus will produce sensory impairment of the contralateral face.

In thalamic disorders, there is often accompanying spontaneous pain or discomfort. These thalamic pains are often very intense and occur in paroxysms affecting the opposite side of the body. They are typically more pronounced in the face, hand or lower leg, and are uncommon on the trunk. Causes include tumours and vascular lesions.

**CORTICAL DISORDERS**

Circumscribed lesions of the postcentral gyros will be followed by localized sensory loss in parts of the opposite half of the body. This is typically a loss of discriminatory sensory function, and includes loss of position sense, impaired ability to localize touch and pain stimuli, elevation of the two-point threshold, and astereognosis. In acute lesions of the parietal cortex, there may appear to be impairment of pain sensibility, but this is rarely prominent or persistent.

Occasionally, sensory seizures are seen in lesions of the sensory cortex, although these are rare. Typically, they show themselves as a wave of sensory irritative symptoms spreading over the body in accordance with the somatotopic organization of the sensory cortex.

**SIMULATED SENSORY LOSS**

A variety of sensory disorders may be simulated. Patients may complain of the sensory loss, but more commonly it is found incidentally during examination. The area of sensory loss is usually sharply demarcated and, in other than medical personnel, does not
conform to a recognized anatomical distribution. Characteristically, pain loss is the most striking feature, and loss of position sense is uncommon. Bearing in mind the difficulties of the sensory examination, it is probably safer to ignore sensory findings that do not fit with any other neurological abnormalities that are present. Often it is possible to make a positive diagnosis of a simulated sensory loss because the individual’s knowledge of anatomy is not sufficient to enable them to get it right. For example, anybody who is found to have unilateral loss of vibration sense on the skull can be confidently diagnosed as simulating. It must be added, however, that the positive finding of non-organic sensory loss does not necessarily indicate that the whole of the neurological problem is simulated. Sensory loss that is simulated is often added on to symptoms and signs as part of a functional overlay.

SEXUAL DYSFUNCTION
Mark Kinirons

Most commonly, advice is sought for problems arising in the context of heterosexual relationships. For women, the two main categories of complaint are disorders of arousal and disorders of orgasm during sexual intercourse. For men, complaints can be grouped as erectile impotence (when erection cannot be obtained or maintained), premature ejaculation and retarded ejaculation. There appear to be quite marked differences between men and women in their expectations from sexual intercourse, the majority of men complaining of problems with erection or ejaculation, and only a minority seeking help on account of lack of interest or enjoyment of sex. An approach to understanding sexual dysfunction must take account of the social, emotional, psychological and physiological effects that may all influence the sexual response, so that, when advice is sought for sexual problems from patients with multiple sclerosis, hypertension, epilepsy, colostomy, mastectomy and many other physical conditions, the clinician must consider the effects of the reaction of the patient and their partner to the disability as well as the direct physiological disturbances and the effects of medication.

Disorders of arousal and orgasm in both men and women are most commonly connected with psychological and emotional factors. Particularly in the young, there may be anxiety about sexual prowess and performance. Negative feelings about intercourse may arise in people with obsessional personalities because of a general fastidiousness and a fear of losing control of feelings and bodily functions. Fear of pregnancy is possibly the most widespread and realistic inhibiting factor. It is also normal for many women to achieve orgasm only through sexual activities other than full intercourse. In women, pain during intercourse (dyspareunia) may result from vaginal infection or other pathology in the pelvis. There may also be inadequate lubrication of the vagina, particularly in postmenopausal or lactating women. In some women, there is a tendency to develop spasm of the pelvic floor muscles during intercourse, resulting in vaginismus, which makes intercourse painful or impossible.

Failure to achieve orgasm (anorgasmia) may be the outcome of dyspareunia or occasionally severe physical malformation of the genital tract, but in the majority of cases there appears to be no good physical or psychological reason for their inability to achieve orgasm. Women depend on tactile stimulation for arousal, and anorgasmia may be due simply to poor technique by the partner.

Libido may be severely impaired in depressive illness. Chronic alcoholism may cause reduced libido in both sexes. Organic causes of impotence include autonomic neuropathy (e.g. diabetes mellitus), spinal cord damage and sometimes central nervous system impairment, as in tumours in the region of the third ventricle and temporal lobe epilepsy. Endocrine disorders include primary and secondary hypogonadism. The only drugs which clearly and directly interfere with vascular mechanisms leading to erection are the ganglion blockers. Adrenergic blockers interfere with ejaculation, and drugs may interfere with the male sexual function by effects on the central nervous system or on blood pressure, or by endocrine effects. For example, the reduced libido in some patients with temporal lobe epilepsy has been attributed to the effect of anticonvulsant drugs in lowering testosterone levels.

Medical advice may be sought by homosexual individuals because of sexual and relationship difficulties, and these may be exacerbated by the attitudes of society. A group of sexual disorders that may lead the person into trouble with their partner or with the law are the disorders of sexual preferences (the paraphilias), which include abnormalities of the sexual object (fetishism or paedophilia) and abnormalities of the sexual act (exhibitionism, voyeurism, frotteurism and sadomasochistic sexual practices). Fetishism and sadomasochism may be quite common and easily concealed in a stable relationship. Help may be
sought only when such practices cause strain in a heterosexual or homosexual partnership. Fetishism occurs almost only in men, and describes the use of objects such as clothing, parts of the body or specific textures such as rubber, leather or plastic to stimulate erection. Sadomasochism related to sexual arousal occurs in both men and women, and has a role more frequently in fantasy than in reality. The sadist inflicts pain, and the masochist adopts a passive role, as in sexual practices involving bondage. Generally, these practices do not lead to conflict with society. A medical opinion is most likely to be sought with sexual offenders, the common deviations being exhibitionism and paedophilia. Exhibitionism or indecent exposure is the gaining of sexual stimulation by exposing the genitalia. Unlike other forms of sexual deviation, there is no attempt to establish further contact with the ‘victim’. In some instances, the exhibitionist may derive satisfaction in a sadomasochistic way by evoking fear in a witness. Some become highly sexually aroused and masturbate during or shortly after the exposure. Frotteurism describes the achievement of sexual arousal by rubbing against a stranger, often in a crowd.

Paedophilia describes sexual activity with children in preference to adults. Usually, a paedophile will be attracted either to boys or to girls, but not to both. It is recognized almost exclusively in men.

Factors sometimes associated with sexual deviations include disturbed rearing and parental disharmony leading to difficulties in establishing stable adult relationships, and personality disorders, particularly in those who are inadequate or psychopathic. The role of low IQ, alcohol abuse and mental illness (manic depressive illness and schizophrenia) should also be considered in the clinical assessment of a case.

Problems with gender identity reflected in cross-dressing behaviour (a number of quite distinct sexual problems. For some men, the wearing of female clothes is sexually arousing, and the clothes are objects of fetishism. A homosexual of either sex may cross-dress out of preference but in the absence of any genuine gender confusion or desire to be of the opposite sex. Finally, there are genuine transsexuals of either sex who have often since early childhood been aware that their psychological sex is opposite to their anatomical sex. These transsexuals may proceed to successful surgical sex reassignment.

**SKIN HARDENING**

**Barry Monk**

Hardening of the skin most commonly arises through the abnormal deposition of collagen in the dermis and subcutaneous tissues, and is the principal feature of *scleroderma*. This occurs in localized and progressive systemic forms. Localized scleroderma, or *morphoea*, usually begins in patients under the age of 30 years as asymptomatic, round or oval, firm, smooth, reddish plaques many centimetres in diameter. They are most common on the trunk, and they may have a lilac or telangiectatic border. After several years, the centre of lesions may become atrophic and hyperpigmented. More rarely, morphoea appears in linear form either in a paramedian distribution on the scalp (*coup de sabre*) or along the line of a limb. Sclerosis in linear morphoea is very marked, and there can be associated atrophy of underlying bone or muscle. The linear type shows little tendency to spontaneous resolution. At times, the whole of a limb or one side of the head may be affected in a child, resulting in a deforming hemiatrophy. Even more rarely, very large areas of skin may be involved in a *generalized morphoea*.

The progressive type of scleroderma is a systemic disease, probably of autoimmune aetiology. It occurs in two distinct forms. In the calcinosis, Raynaud’s phenomenon, sclerodactyly, telangiectases (CRST) syndrome, symptoms of Raynaud’s phenomenon are followed by progressive *acrosclerosis* (hardening of the extremities), with an associated loss of mobility; calcinosis of the fingertips may be a prominent feature. In addition, telangiectasia is seen, and the skin on the face – especially around the mouth – becomes tight and fixed. Disorders of oesophageal mobility are common, and pulmonary hypertension is a late, but ominous, feature (Figs S.22 and S.23).

In the more generalized form of progressive systemic sclerosis, the whole skin becomes thickened and immobile; the lungs, kidneys and heart may also be affected, and there is little tendency to spontaneous improvement. Features mimicking the hardening of the skin in scleroderma may be seen occasionally in the late stages of *porphyria cutanea tarda*, in *carcinoid syndrome*, and in patients affected by occupational exposure to the manufacture of vinyl chloride.

Hard, bead-like infiltration of the skin is seen with *lipoid proteinosis*, and this is particularly noticeable around the eyelid margins. Hardening of the skin of the chest can follow secondary infiltration with *scirrhous carcinoma*. The skin of the lower legs may...
become hard in patients with long-standing venous insufficiency (lipodermatosclerosis), and a general hardening of the skin may be evident in diabetes of long standing.

GENERALIZED HYPERPIGMENTATION

Diffuse hyperpigmentation, including pigmentation of the buccal mucosae, gingival margins, palmar creases and nailbeds, is a feature of Addison’s disease. In fact, this disorder is remarkably rare in clinical practice, and a more likely cause is the presence of an ectopic hormone-secreting tumour; for example, various lung carcinomas secrete secrete peptides, some of which have melanocyte-stimulating hormone-like properties. Similarly, adrenocorticotrophic hormone for intramuscular injection is often contaminated with similar peptides and can cause hyperpigmentation. Diffuse pigmentation can be a feature of malignant cachexia, the late wasting phase of HIV disease, and can be very marked in those dying with a heavy tumour load of secondary malignant melanoma, sometimes accompanied by melanuria. The diffuse hyperpigmentation of haemochromatosis (‘bronze diabetes’) is due to an excess of both melanin and iron.

LOCALIZED HYPERPIGMENTATION

Many skin disorders appear able to cause a non-specific post-inflammatory hyperpigmentation apparently related to leakage of melanin from its normal location. This is particularly seen following lichen planus (Fig. S.24) and in a fixed drug eruption, where the condition can be diagnosed retrospectively from the distribution of the pigmentation. Dark-skinned races seem more susceptible to the phenomenon of post-inflammatory changes in pigment, and in lichen planus in South Asian races, for example, this can be most disfiguring. In some chronic conditions, pigmentation forms part of the spectrum of diagnostic physical signs, for example, atopic dermatitis, morphea, urticaria pigmentosa and acanthosis nigricans. In a berloque dermatitis, an acute phototoxic reaction occurs to perfume or eau de cologne, due to the psoralsens that they contain. In its most florid form, the condition can be dramatic and even bullous, but patches of hyperpigmentation on the sides of women’s necks are extremely common. A similar hyperpigmentation can follow a photo-toxic reaction to plant chemicals – a photophytodermatitis – as seen with plants of the Umbelliferae family, such as wild parsley and parsnip.
A common cause of a symmetrical hyperpigmentation on the forehead and cheeks in middle-aged female subjects is chloasma (melasma). This condition is of unknown cause, but may be exacerbated by pregnancy or the oral contraceptive pill, and by sunlight. Less common causes of localized hyperpigmentation include lichen amyloid (Fig. S.25) and naevus of Ota (Fig. S.26).

HYPERPIGMENTATION OF EXPOSED AREAS

Ultraviolet (sunlight) exposure will tan the skin in those with appropriate skin types, but tanning may be accentuated by certain drugs, including thiazides, sulphonamides, amiodarone, nalidixic acid, tetracyclines and chlorpromazine. A similar phenomenon may arise in metabolic disorders such as pellagra and porphyria cutanea tarda.

HAEMOSIDERIN

Haemosiderin is a breakdown product of haemoglobin, and will be precipitated in the skin, leaving a persistent, localized, orangey-brown discoloration in situations where there has been chronic extravasation of blood, as in stasis ulceration of the legs, and in some cutaneous vasculitic disorders (pigmented purpuric eruption and Majocchi–Schamberg disease) (Fig. S.27).

OTHER CHEMICALS

Obstructive liver disease causes acute or insidious jaundice of differing hue, for example the greenish colour of primary biliary cirrhosis. Carotinaemia results in a yellowish or orange discoloration of the skin, sparing the sclerae, and follows the ingestion of large quantities of carrots, oranges and other vegetables. Yellowing of the skin can be caused by drugs, particularly mepacrine. In alcaptonuric ochronosis, the pigment precursor tyrosine cannot be catabolized, and a bluish-black pigment accumulates, particularly in the cartilaginous tissues.
of the ears, sclerae, joints and vertebral column. The urine becomes black on standing for 24 hours. Topical use of hydroxyquinone-based chemicals, which are sold in some countries as skin lighteners, can also produce a permanent deep pigmentation due to ochronosis. The history in such cases may be misleading, as patients may be too embarrassed to admit use of such products, but the histology is characteristic. A generalized bluish discolouration is seen with methaemoglobininaemia, which can be drug induced (e.g. by dapsone). A deeper colour is induced by the deposition of silver in argyria; this may arise from chronic occupational exposure or from the medicinal use of silver salts. High-dose, long-term treatment of acne or rosacea with minocycline can result in pigment deposition within acne scars, and sometimes more widely (Fig. S.28). Zidovudine (AZT) can cause linear pigmentation of the nails (melanonychia striata) and sometimes diffuse hyperpigmentation.

Tattooing of the skin is usually obvious, but may produce unusual patterns when it has arisen accidentally from blast injuries (Fig. S.29). Occasionally, bizarre colouring of the skin arises from attempts to fool doctors or alarm relatives (factitious dermatitis).

**SKIN TUMOURS**

Barry Monk

Cutaneous tumours are solid lesions of the skin that are larger than nodules (conventionally greater than 1 cm). They must be distinguished from cysts, although a necrotic tumour may form a cystic centre. Tumours may arise in benign and malignant growths, but they may also be seen in inflammatory disorders as diverse as leprosy and sarcoid.

A classification of skin tumours is listed in Box S.5.

**MALIGNANT TUMOURS**

Secondary deposits

These commonly occur late, as multiple, hard, fleshy nodules of any colour (Fig. S.30). Subcutaneous secondaries can masquerade as benign lesions, and a high level of suspicion should be maintained. Certain areas are predisposed to metastases, including the scalp (breast, lung and genitourinary tract carcinoma),
the chest wall (breast cancer) and the abdominal wall, especially around the umbilicus (carcinoma of the stomach and the colon). The cell type identified on histology may give a clue to the primary source. The skin is sometimes infiltrated by leukaemia or lymphoma, the patients presenting with multiple, randomly scattered, erythematous nodules; it is a late, and usually pre-terminal, manifestation. Paget's disease (Fig. S.31) is invariably associated with an underlying malignancy of the breast.

Kaposi's sarcoma
This is a malignant proliferation of vascular endothelial cells giving rise to superficial, subcutaneous or deeper vascular tumours (Fig. S.32). The tumours do not metastasize, but are multi-site in origin. They are probably due to infection with human herpes virus-8. Tumours flourish when immunity is depressed, for example in HIV, post-transplant and in old age. The classical lesions, which were first described in patients of Jewish or Italian descent living in Vienna in the 1870s, are probably acquired on the genome and expressed in old age. Endemic Kaposi's sarcoma was first recognized in the 1950s as a common tumour in sub-Saharan Africa, in younger patients, and was seen to follow a much more aggressive course.

Individual lesions begin as pink vascular macules, often in multiple sites, and they gradually enlarge and become palpable and darken with time. Draining oedema is often prominent. The vascular lesions can simulate granulomas, histiocytomas or haemangiomas. Internal lesions can occur chiefly in the gastrointestinal tract and lungs. If immunity improves (i.e. with antiretroviral therapy), the tumours may shrink.

Primary malignant skin tumours

Squamous cell carcinoma (Fig. S.33) is usually single and, as a rule, is fairly slow-growing, extending peripherally and infiltrating deeply while ulcerating at its centre. Eventually, the lymphatic nodes draining the affected area become involved and enlarged. The usual sites for squamous cell carcinoma are the lips, especially the lower lip and sun-exposed areas, as well as glans penis and vulva. Solar keratoses, X-ray scars and lupus vulgaris may all undergo malignant squamous change. The main diagnostic features are its origin as a single growth, its craggy hardness, its slow development, and the metastases to neighbouring lymph nodes.
Malignant melanoma (Fig. S.34) can arise anywhere on the skin surface at any age, although it is rare in childhood; it may arise de novo or from a pre-existing mole. It is rare in black individuals and increases in frequency with the fairness of the skin and the amount of previous sun exposure. Patients with a history of severe sunburn in childhood, with more than 50 moles on their skin, more than five unusually large moles or a family history of malignant melanoma are at increased risk. This tumour may occur on the scalp, under a nail or on anogenital skin.

Occasionally, melanomas lose the capacity to produce pigment; this is termed amelanotic melanoma. The characteristics of malignant melanoma are its rapid development and growth, its deepening colour, its ulceration, areas of depigmentation, bleeding and crust formation, and its rapid metastases. Sometimes multiple metastases occur in the skin itself. The disease is highly malignant, but early diagnosis and excision is curative. The prognosis depends on the thickness of the primary lesion at the time of excision.

Rodent ulcer (basal cell carcinoma) (Fig. S.35) usually affects the face (see FACE, ULCERATION OF, p. 188). These are the most common primary skin malignancies. They do not metastasize, but local invasive destruction can be extensive if the lesions are neglected.

Mycosis fungoides (cutaneous T-cell lymphoma) (Fig. S.36) is a rare, chronic, slowly fatal disease that is characterized in its final stage by tomato-like growths that may ulcerate. For many years, a 'pre-mycotic', non-specific, red scaly rash is present, later (sometimes over 30 years) forming red plaques of differing hue, and finally tumours.

Xeroderma pigmentosum is an extremely rare disorder of nuclear protein repair that is inherited as
a recessive trait. It presents in childhood as a proneness to sunburn and early gross sun damage with elastosis, atrophy, telangiectasia and finally multiple skin tumours (squamous cell carcinoma, rodent ulcer, malignant melanoma and kerato-acanthoma).

**BENIGN CUTANEOUS TUMOURS**

**Lipomas** are usually multiple subcutaneous nodules, which may be lobulated and found on any part of the body. They occur as a familial trait but can often not be discerned until adulthood. **Seborrhoeic keratosis** (basal cell papillomas) (Fig. S.37) are extremely common and start as papules in middle age, growing into larger, flat, greasy, warty, pigmented neoplasms. They are sometimes unkindly called ‘senile keratoses’.

A **cutaneous horn** is a peculiar cutaneous neoplasm surmounted by a spectacular horny overgrowth. The nature of the underlying neoplasm can only be safely diagnosed by examining the histopathology, for example actinic keratosis, squamous cell carcinoma, viral wart, kerato-acanthoma and seborrhoeic wart (basal cell papilloma).

A **pyogenic granuloma** (granuloma telangiectaticum) is a fairly common skin lesion. It often develops at the site of a recent injury and is composed of proliferating capillaries in a loose stroma. This produces a rapidly growing vascular nodule that bleeds easily when traumatized. It is distinctive, being bright red, 0.5–1 cm in diameter, often pedunculated, and surrounded by a collar of thickened epidermis. The most common sites are the fingers, upper chest, lips and toes. It must be differentiated from amelanotic melanoma and glioblast tumour. Kaposi’s sarcoma in HIV-infected patients can accurately mimic pyogenic granuloma, and for this reason histological analysis after curettage or excision is mandatory.

The superficial dermis contains many specialized tissues, some of ectodermal and some of mesodermal origin, forming the various adnexal structures (e.g. hair, sebaceous glands and sweat glands). Benign, or rarely malignant, neoplasms of all these specialized tissues can occur, for example leiomyoma, hyaladenedoma, neurofibroma, sebaceous adenoma, tricho-epithelioma and glioma tumour. The diagnosis is often first made by the pathologist.

Tumours may be seen in a variety of inflammatory disorders. Rheumatoid nodules are mobile and arise over the extensor surfaces of the knees and elbows. Sarcoïdosis of the skin may present with papules, nodules or tumours, or with extensive plaques, often with a rather mauvish hue. Evidence of sarcoid in other tissues may suggest the diagnosis. Tumorous lesions of the skin may be seen in lepromatous leprosy in those living in endemic areas.

**SLEEP, DISORDERS OF**

Mark Kinirons

The most common complaint is of insomnia, although some patients will seek advice on account of too much sleep (hypersomnia) or occasionally for abnormal events that occur during sleep (the parasomnias) (Box S.6).
lacks a daytime schedule and has no motivation to dreading the coming day. 2–3 hours earlier than normal, feeling depressed and or melancholic depression. Such subjects may wake awakening being one of the features of endogenous initial, middle and late insomnia, early morning of worrying thoughts. In depression, there may be show little sign of fatigue, whereas the anxiety-prone personalities. Such people tend to be highly aroused under stress at home or at work, or who have anxious complaints of poor sleep come from people who are under stress at home or at work, or who have anxious personalities. Such people tend to be highly aroused and stay awake thinking about stressful situations and planning how to cope. In others, their reports of poor sleep may be due to stimulants such as coffee, tea or smoking cigarettes, and there is a high rate of reported poor sleep in patients with an alcohol problem. The normal requirement for sleep varies widely, and a small number of individuals require only 3–4 hours of sleep each night. During a painful physical illness, or in a person with respiratory difficulties, the causes of disturbed sleep will be obvious. Poor nutritional status and low weight may also be accompanied by diminished sleep.

A change in sleep pattern is extremely common in all forms of mood disturbance. The manic patient may remain cheerful and active throughout the night and show little sign of fatigue, whereas the anxiety-prone patient may have difficulty falling asleep because of worrying thoughts. In depression, there may be initial, middle and late insomnia, early morning waking being one of the features of endogenous or melancholic depression. Such subjects may wake 2–3 hours earlier than normal, feeling depressed and dreading the coming day.

Hypersomnia may be the complaint of someone who lacks a daytime schedule and has no motivation to rise. Poor motivation and the effects of medication can account for the apparently increased sleep of some psychiatric patients, especially those with schizophrenia.

In narcolepsy, the person experiences bouts of drowsiness leading to short periods of sleep of a few minutes’ duration recurring two or three times a day. The condition may be associated with cataplexy, a sudden loss of muscle tone lasting for a few seconds, often triggered by strong emotions. Other features of the condition include sleep paralysis, in which the subject is momentarily paralysed and unable to move (as happens sometimes with normal people awaking from a bad dream), and hypnagogic hallucinations, which may be auditory, visual or tactile. Usually, the diagnosis of narcolepsy is made from the clinical history and electroencephalography findings that on falling asleep the patient goes spontaneously into rapid eye movement (REM) sleep, without passing through a non-REM stage.

One rare form of hypersomnia that occurs usually in young men is the Kleine–Levin syndrome. The patient sleeps excessively by day and night but is rousable. Such patients often eat excessively. Organic lesions of the midbrain or hypothalamus may also cause increased hunger, weight gain and drowsiness. The abrupt onset of daytime sleepiness or drowsiness should, however, immediately alert the clinician to the possibility of an intracranial space-occupying lesion. When unusual behaviour patterns during sleep develop suddenly in a patient, a drug effect should be rapidly excluded.

Nightmares are a normal phenomenon with no psychiatric significance. Frequent nightmares may occur during anxiety states and depression, in post-traumatic stress disorders, and with alcohol abuse or following a change of hypnotic. Sleep-walking and night terrors may be familial, and can be precipitated by drugs such as antidepressants, anticonvulsants, analgesics, lithium and phenothiazines. These sometimes follow a febrile illness. A child experiencing a night terror that usually occurs within the first 2 hours of sleep may sit up with an expression of fear and remain oblivious to the surroundings for a few minutes before dropping soundly to sleep again. The child will have no memory of the event in the morning. Sometimes, sleep-walking occurs during a night terror. Usually, patients will sleep-walk in a calm, gentle way, but sleep-walkers are at high risk of injuring themselves by falling downstairs or through windows. The subject is in a state of automatism and may walk some distance and carry out quite complicated series
of actions. Both night terrors and sleep-walking are more frequent during times of stress, and families will require reassurance.

In bruxism, the grinding of teeth during the night causes dental problems and can be provoked by psychotropic medication. Nocturnal enuresis is a common occurrence, and multifactorial sources of stress should be sought in a family when this develops in a child after a period of established bladder control.

SLEEP APNOEA/HYPOPNOEA SYNDROME

The major complaint of patients with the sleep apnoea/hypopnoea syndrome is usually daytime sleepiness, which may vary in severity from trivial to dangerous. Indeed, many patients fall asleep during driving, working or in mid-conversation. The patient is usually unaware of the hundreds of brief awakenings per night that result in the daytime sleepiness, although around one-third of patients are aware of awaking occasionally at night with choking episodes. The sleep disruption results from total or partial occlusion of the upper airway at the level of the soft palate or tongue each time a patient goes to sleep. This produces apnoeas or severe hypopnoeas that are only terminated when the patient awakens briefly, perhaps due to the fall in arterial oxygen levels. The awakening is so brief that the patient is not aware of it, but it is sufficient to increase the tone to the upper airway-opening muscles, and breathing resumes for a few seconds until the patient goes back to sleep and becomes apnoeic or hypopnoeic again. Patients do not find nocturnal sleep satisfying, and they sometimes awaken with a headache. Their bed partners report very loud snoring punctuated with breathing pauses, and that the patient is a very restless sleeper, often thrashing around the bed.

This condition affects around 1 per cent of adults, with 85 per cent of patients being male and 50 per cent being overweight. Severe long-standing cases, particularly in patients with co-existing lung disease, may be complicated by cyanosis (see CYANOSIS, p. 120) and right heart failure.

SMELL, ABNORMALITIES OF

Michael Gleeson

Abnormalities of the sense of smell are commonplace, and most people will have experienced this at some time or other. Although some claim increased sensitivity to odours (hyperosmia), this is not a pathological condition, merely an expression of normal variation. Similarly, some normal people seem perversely unaware of smells (hyposmia) yet are able to perceive other odours. This too is rarely abnormal. Complete loss of the sense of smell (anosmia) through one or both nostrils is of importance as it may be treatable and can be a sign of potentially serious disease. Perhaps more disturbing for the patient is distortion of the sense of smell (dysosmia), or the perception of non-existent odours (parosmia). The common causes of anosmia are listed in Box S.7.

Reduced or total loss of the sense of smell is a common accompaniment of rhinitis and rhino-sinusitis. Polyps are usually clearly visible by inspection of the nasal cavities, and occasionally in severe cases protrude from the nose (Figs S.38 and S.39). Sometimes, mucopus can be seen escaping from under the middle turbinate, indicating paranasal sinus infection. However, signs of infection may not be apparent on clinical examination and are only evident after scanning the patient.

<table>
<thead>
<tr>
<th>Box S.7 Causes of anosmia</th>
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<tr>
<td>Nasal obstruction</td>
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<tr>
<td>• Rhinitis and rhinosinusitis</td>
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<tr>
<td>• Deviated nasal septum</td>
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<tr>
<td>• Sino-nasal malignancy</td>
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<tr>
<td>Abnormalities of the olfactory mucosa</td>
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<tr>
<td>• Rhinitis medicamentosa and atrophic rhinitis</td>
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<td>• Post-viral neuropathy</td>
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<tr>
<td>Abnormalities of the olfactory nerves and pathways</td>
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<td>• Congenital absence</td>
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<tr>
<td>• Trauma</td>
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<tr>
<td>• Neoplasia (olfactory neuroblastoma, meningioma)</td>
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<tr>
<td>• Alzheimer’s disease, Parkinson’s disease, Huntington’s chorea, temporal lobe epilepsy</td>
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<tr>
<td>Psychological</td>
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Figure S.38 A simple nasal polyp that was easily seen by elevation of the nasal tip.
Treatment of rhinitis or rhinosinusitis by medical or surgical means may restore the sense of smell for some. Patients who lose their sense of smell after viral infection, but have no obvious residual sepsis, may also be helped by topical steroid drops, but these fortunate patients are few and far between. Those who lose their sense of smell as a result of an anterior cranial base fracture rarely, if ever, recover it.

Tumours affecting or arising in or around the cribriform plate of the ethmoid produce a slowly progressive deterioration and loss of the sense of smell. Olfactory neuroblastoma (aesthesioneuroblastoma), meningioma and juvenile angiofibroma are the most common tumours in this region, and it is prudent to remember that they may develop in relatively young patients. Loss of the sense of smell is an early symptom and sign in these patients, and it becomes established well before proptosis, nasal obstruction and blood-stained nasal discharge (Fig. S.40).

Parosmia is a perverted sense of smell that is characteristically unpleasant and usually develops after damage to the olfactory nerves. Common causes include head injury, tumours and nasal infection. Olfactory hallucinations represent a common and striking symptom of partial seizures originating in the medial aspect of the temporal lobe (uncinate seizures). These hallucinations tend to be pungent, sometimes likened to burning rubber or a smell of gas. Olfactory hallucination is not uncommon in psychosis, when unpleasant smells may be attributed either to the patient him-or herself, or to others. Patients with rhinoliths sometimes describe an offensive smell that is not always apparent to others.

SNEEZING

Michael Gleeson

Sneezing is usually initiated by irritation of the nasal mucosa, especially that of the anterior part of the nasal septum or the turbinates. The sneeze reflex is protective in nature and essentially similar to the cough reflex. It is a sudden and involuntary expulsion of air through the nose and mouth, and it is controlled by a reflex initiated by the Vth and Xth cranial nerves. Sneezing may be caused by local irritation or increased sensitivity of the nasal mucosa. The early stage of the common cold is probably the most common cause of sneezing, which subsides once the bacterial secondary infection supervenes with a thicker nasal discharge. Sneezing is common in the prodromal stages of measles and the other infectious fevers, but it is not common in more chronic nasal infections.

The inhalation of fine dusts and powders causes sneezing due to local irritation. Riot control gas and war gases irritate the eyes as well as the nasal mucosa; lacrimation, nasal discharge and sneezing may be incapacitating to those exposed. Some individuals are particularly sensitive to changes of temperature and light. Many people sneeze once or twice on their first exposure to bright light. Bouts of sneezing are frequently observed in individuals with atopic sensitization to airborne allergens. When the nasal mucosa encounters an allergen to which the individual is sensitized, inflammatory mediators are released from mast cells and basophils, and these may cause sneezing.
SNORING
Michael Gleeson

Snoring is the noise produced in sleep by the vibration of the soft palate and the tongue base. A differentiation is made between simple snorers and patients with obstructive sleep apnoea (OSA) (see also SLEEP, DISORDERS OF p. 623). Whereas in simple snorers the blood oxygen saturation does not fall significantly, in patients with OSA the upper airway collapse results in recurrent periods of apnoea, blood oxygen desaturation and subsequent arousal reactions. This disorder may lead to fatigue, loss of energy during the day, changes in personality and, in severe cases, pulmonary hypertension and right ventricular strain.

It is now considered that a continuous spectrum exists between uncomplicated simple snoring and the OSA syndrome. Simple snorers may develop OSA after the ingestion of alcohol. Snoring is more common in obese people, and in many patients a change in lifestyle (loss of weight and reduction of alcohol consumption) may improve their symptoms. In selected cases, appropriate surgical treatment aiming at the portion of the airways that is responsible for the vibration or collapse may be beneficial. In children, snoring is most often due to adenoid or tonsillar hypertrophy, which may be cured by adenotonsillectomy. Patients with nasal obstruction may benefit from septoplasty or sinus surgery. If redundancy of the soft palate and lateral pharyngeal walls are the source of the problem, surgical resection may be effective, a procedure known as uvulopharyngopalatoplasty. Cases of tongue base collapse are more difficult to treat surgically, but this may be improved by use of a mandibular advancement device.

SPEECH, ABNORMALITIES OF
David Werring & Mark Kinirons

Speech and language are complex functions that are critical for normal human social interaction. Language disorders involve the abnormal production or comprehension of written or spoken language; those due to acquired brain disease (usually focal) are termed dysphasia or aphasia. Although speech production is often disrupted in disorders of language, this entry is limited to speech disorders in which language function is preserved. The disorders to be considered here are dysarthria and dysphonia.

Dysarthria is a motor disorder of speech production due to dysfunction of the muscles of articulation. Dysphonia is a motor disorder of voice production.

Appraisal of speech is a rarer disorder of speech motor function unaccounted for by disease of the muscles of articulation or their innervation, and is considered elsewhere (see APRAXIA, p. 33).

A classification of developmental speech and language delay is provided in Box S.8.

DYSARTHRIA

Speech production requires a complex integration of motor actions, and consists of respiration, phonation, articulation, resonance and prosody. The respiratory muscles are important so that expiration occurs with appropriate duration, force and flexibility. Intact anatomy and function of the respiratory muscles, larynx, palate, nasal passages, tongue and lips are required for normal speech. The respiratory muscles (intercostal muscles and diaphragm) are supplied by the phrenic nerve, which arises from cervical segments 3, 4 and 5. The larynx, pharynx, tongue and lips receive lower motor neurone innervation from the trigeminal, facial, vagus and hypoglossal cranial nerve nuclei. The nuclei of these nerves are in turn controlled by upper motor neurone descending pathways from the speech motor cortex (the corticobulbar tracts). Pure dysarthria does not involve a dysfunction of cortical language functions, so that comprehension, reading and writing are intact. It is convenient to subdivide dysarthria into lower motor neurone, upper motor neurone (pseudobulbar), cerebellar (ataxic), hyperkinetic and hypokinetic. The dysarthria that commonly results from a unilateral cortical or subcortical stroke does not fall into any of the above categories.

Lower motor neurone dysarthria

Involvement of the lower motor neurone innervation of the bulbar muscles (cranial nerve nuclei or below) produces this type of speech, which is characterized by weak, imprecise articulation with poor breathing support. Air escaping through the nose due to palatal weakness causes hypernasality. The tongue may be weak and fasciculating, palatal movements may be reduced, and the ability to cough explosively may be lost due to vocal cord paralysis. This pattern may
also termed a bulbar palsy. The lesion can be in the cranial nerve nuclei, the cranial tongue. Causes include the motor neurone diseases, although in the most common form – amyotrophic lateral sclerosis – upper motor neurone features will also be present. Pure lower motor neurone diseases include X-linked bulbospinal neuronopathy (Kennedy syndrome, with gynaecostasia and diabetes), multifocal motor neuropathy and the post-polio syndrome. Other causes of bulbar palsy include cranial polyneuropathies due to meningeal disease, myasthenia gravis, lower brainstem stroke, or muscle diseases involving the bulbar muscles such as polymyositis, dermatomyositis or oculopharyngeal muscular dystrophy.

Upper motor neurone dysarthria
Involvement of the upper motor neurone pathway to the bulbar cranial nerves produces a pseudobulbar palsy. The type of speech that results is termed spastic dysarthria; the tongue is immobile and spastic, so that speech has a strangled, effortful quality, sometimes with associated spastic dysphonia, and it is often slow. Bilateral involvement of either the cortex or descending pathways is necessary to produce pseudobulbar palsy although, in the case of multiple strokes, there may be old pre-existing lesions followed by a strategically placed acute infarct causing a sudden presentation with dysarthria or anarthria (an inability to produce any intelligible speech). Other causes of pseudobulbar palsy include motor neurone disease with upper motor neurone involvement, multiple sclerosis, large frontal tumours and cerebral palsy. Associated features of a classical pseudobulbar palsy (reflecting damage to frontal cortex descending projections) are emotional lability with inability to suppress laughter or crying, and a ‘frontal’ gait disturbance with small stuttering steps, termed marche à petit pas.

Cerebellar dysarthria
Cerebellar dysarthria may present with slurred speech, superficially resembling alcohol intoxication. However, there are often additional problems of ‘scanning’ or explosive speech, due to an inability to modulate accurately the rhythm, rate and force of speech. Cerebellar dysarthria may result from focal damage to the vermis or more widespread damage to the whole cerebellum (or its connections). Any disease affecting the cerebellum can cause this pattern of dysarthria; causes include multiple sclerosis, cerebellar degeneration (e.g. paraneoplastic cerebellar degeneration, spinocerebellar degenerations, alcohol), stroke, vitamin E deficiency, hypothyroidism and coeliac disease.

Extrapyramidal dysarthria
Extrapyramidal diseases result from pathology of the basal ganglia, are attributed to neurotransmitter abnormalities (e.g. dopaminergic neurone loss in Parkinson’s disease), and cause abnormal control of volitional movement. Some extrapyramidal diseases (e.g. Parkinson’s disease) cause slowness of movement and rigidity (hence the term ‘akinetic–rigid syndrome’), which may impact on speech as well as limb motor control. In other extrapyramidal disorders (e.g. Huntington’s disease), basal ganglion pathology causes an excess of involuntary movements that interfere with normal volitional movement. It follows from the above that two types of speech disorder may be seen in extrapyramidal diseases: hypokinetic and hyperkinetic. In Parkinson’s disease, the speech is hypokinetic. The range of articulation is reduced, breathing support is poor, and the volume is low. The speech may be either slow and monotonous or produced in short, rapid phrases. In the early stages, the only abnormality might be a slight loss of prosody (emphasis and inflections); in the late stages of Parkinson’s disease, the speech may be virtually inaudible and unintelligible. Palilalia may be seen, in which there is repetition of a phrase, which the patient retries with increasing rapidity. Hyperkinetic dysarthria results from chorea, orofacial dyskinesia and athetosis (see CHOREA, p. 100). Articulation is interrupted by involuntary movements, producing jerky and irregular speech with reduced intelligibility.

DYSPHONIA
Spastic dysphonia
Spastic dysphonia, also termed spasmodic dysphonia or laryngeal dystonia, is a form of focal dystonia (i.e. a disorder of involuntary muscle movement and postures), thought to be due to abnormal functioning in the basal ganglia. It may be primary or secondary to another neurological disorder. The muscles of the larynx contract so that, in the adductor type, the vocal cords are brought together and speech is effortful, strained and strangled. Symptoms may improve or disappear temporarily, either spontaneously or when yawning, laughing, singing or relaxing. In the abductor type, there is an excessive action of the muscles that open up the vocal cords, resulting in a disjointed, breathy, whispering voice pattern. Adductor and abductor spasms may co-exist, causing an irregular voice tremor. Upper motor neurone disorders affecting the vagus – and hence the muscles innervated by the recurrent laryngeal nerve – may produce a spastic dysphonia, which can be associated with spastic dysarthria.
Flaccid dysphonia
Lower motor neurone damage to the vagus nerve, the recurrent laryngeal nerve or the vocal cord muscles may cause the more common flaccid dysphonia. Patients may also have a ‘bovine cough’. This type of dysphonia most commonly results from isolated damage to one of the recurrent laryngeal nerves. It may also be seen in poliomyelitis, motor neurone disease, cranial polyneuropathies, myasthenia gravis and some rare forms of muscle disease.

SPINE, DEFORMITY OF
Fred Heatley and Jonathan Lucas
DEFINITIONS OF TERMS
Unfortunately, these are confusing since the same terms, kyphosis and lordosis, are used to describe both normal and abnormal curves:

- Kyphosis: this is a smooth flexion curve.
  - Normal thoracic kyphosis describes the normal gentle forward curve of the thoracic spine.
  - Postural kyphosis describes a poor posture with ‘drooping shoulders’, which is voluntarily correctable.
  - Compensatory kyphosis means an increased curve of the thoracic spine secondary to some other fixed deformity such as increased lumbar lordosis.
  - Structural kyphosis is fixed and is associated with changes in the shape of several vertebrae with increased wedging anteriorly. Causes include Scheuermann’s disease (adolescent kyphosis), osteoporosis and ankylosing spondylitis.

- Kyphos: this is also referred to as a ‘gibbus’.
In contradistinction to a kyphosis, a kyphos is a sharp, acute-angle deformity due to localized collapse or wedging of one or more vertebrae. A kyphos due to the collapse of just one vertebra is surprisingly easy to overlook. The best way to detect it is to run a finger down the spine with the patient lying prone. A progressive kyphos has a high risk of being complicated by paraplegia. Therefore this deformity is an important sign to be able to detect. The changes on a lateral X-ray are always more dramatic than the physical signs.

- Lordosis: this is a smooth extension curve.
  - Normal cervical or lumbar lordosis describes the standing posture of these two areas of the spine.
  - Compensatory lordosis indicates an increased curve secondary to a structural kyphosis of the thoracic spine.

- An increased structural lordosis does not fully correct on forward flexion and is associated with underlying bony abnormality, for example a spondylolisthesis.
- Scoliosis: this is a lateral curve. It is designated left or right according to the direction of the convexity of the curve. In reality, it is not a simple lateral curve as it is always associated with a rotational deformity of the spine. The bodies rotate towards the convexity, and the spines and neural arches rotate into the concavity.
- Spondylolisthesis: this is a forward slip of a vertebral body on the vertebra immediately below.

Kyphosis
Congenital kyphosis occurs when there is a failure of vertebral segmentation involving several vertebrae and the anterior bone block acts as a tether.
Adolescent kyphosis (Scheuermann’s disease) is, as the name implies, a condition that starts at puberty. The patient becomes progressively more round-shouldered, and there is a smooth thoracic kyphosis with a compensatory lumbar lordosis. Boys are affected twice as commonly as girls. Approximately 50 per cent of children will have some pain over the apex of the curve, but this is usually mild. Neurological complications can occur but are rare. Very occasionally, if the deformity is severe, it may interfere with normal lung function.
Inflammatory kyphosis (also termed ‘ankylosing spondylitis’; see BACK, PAIN IN, p. 50) is a condition that usually progresses from the sacroiliac joints to involve the lumbar spine, thoracic spine and finally the cervical spine. It is usually associated with a kyphotic deformity, a straightening out of the lumbar lordosis and increased thoracic kyphosis. These deformities are often associated with fixed flexion deformities of the hips that greatly exaggerate the bent posture (see Figs B.3 and B.4).

Metabolic causes of kyphosis
Osteoporosis is the most common cause of an increased thoracic kyphosis in which anterior wedging of the bodies of the vertebrae produces a structural deformity. The kyphosis can be so severe as to result in impingement of the costal margin onto the iliac crest. If the deformity is of relatively sudden onset, it is important to exclude other serious underlying diseases such as multiple myeloma. It is also easy to assume that the cause is osteoporosis and to miss the diagnosis of osteomalacia.

Kyphos
Due to the sharp, acute-angle deformity, there is a much greater risk of neurological involvement,
particularly when it affects the thoracic spine. The cord is at greater risk of compression than the cauda equina, which commences at the level of the L1 vertebra.

Congenital kyphos results from a defect of the formation of one or more vertebral bodies. In mild cases, there may be a failure of just part of a single vertebra; in severe types, there may be a total absence of the vertebral body. The deformity increases markedly during growth, and it may lead to paraplegia.

Traumatic kyphos results from a wedge crush fracture. The most common cause is osteoporosis, but the differential diagnosis includes metastatic disease and multiple myeloma. A sharp, angulated kyphos can also occur after a major fracture dislocation, the thoracolumbar region being the most commonly affected site.

Infective kyphos due to pyogenic infection is increasing in frequency due to the longevity of the population in general and the rising incidence of specific illnesses, for example diabetes, HIV and drug addiction. It is an easily missed cause of pyrexia in intensive care units, where it is usually secondary to blood-borne infection from intravenous cannulae. Classically, X-rays show destruction of the intervertebral disc and adjacent endplates – ‘two bodies and the intervening disc’ (see SPINE, TENDERNESS OF, p. 632, and Figs S.51–S.53).

Chronic infection: tuberculosis
Classically, this is a paradiscal infection affecting the anterior distal aspect of the ‘body above’ and the anterior superior aspect of the ‘body below’. While cervical spinal tuberculosis is the most common site in children, thoracic disease classically affects adolescents and young adults. As with pyogenic infection, the lateral X-ray shows the involvement of two bodies and the intervening disc. With progression of the disease, the adjacent vertebral bodies become involved, and the predominantly anterior destruction of the bodies leads to a sharp angular kyphos (Fig. S.41). Large abscesses may develop and migrate considerable distances, for example a psoas abscess presenting in the groin (Fig. S.42). Spinal cord compression (Pott’s paraplegia) may supervene, either with the active disease due to a compression by caseous pus, inflammatory oedema and angulation, or late in the healed disease due to stretching of the cord over the bony ridge of the apex of the deformity. In the former situation, where the compression is ‘soft’, there is usually a good response to conservative therapy, antibiotics and, if necessary, drainage (now possible by aspiration under computed tomography guidance). In the latter situation, surgery should be considered to correct the kyphos to decompress the spinal cord, but is associated with risks of intra-operative cord injury and must be undertaken with intra-operative spinal cord monitoring.

Lordosis
An increased lumbar lordosis is most commonly compensatory, secondary to fixed flexion deformities of the hips, and is therefore found in nearly all hip disease. An increased lumbar lordosis is one of the classical signs of untreated congenital dislocation of the hips and also infantile coax vara.

Scoliosis
There are two fundamental types of scoliosis: postural and structural (Box S.9).

Postural scoliosis
This is a curvature of the spine that corrects on sitting or lying. The commonest cause is a leg length discrepancy. The actual spinal deformity is better described as a tilt or list of the spine as there is no associated vertebral curve or curves.
Structural scoliosis
This is a fixed lateral curvature of the spine associated with vertebral body rotation which is at its greatest at the apex of the curvature. The curve is therefore associated with a rib or loin hump depending on the location of the curve or curves. (Figs S.43–S.45).

Idiopathic scoliosis is the commonest cause of a scoliosis in otherwise outwardly normal children. However, it is very important to undertake a very thorough clinical and neurological examination to exclude other causes for the scoliosis and neural axis imaging is usually indicated and a whole spine MRI scan should be undertaken. The risk of progression of the curve is determined by age and size of the curve at presentation. As growth is the greatest driver of progression of the curve, the younger and more skeletally immature the child presents, the greater the risk of progression. Children need to be monitored to define the rate of progression so intervention can be instigated at the optimal time. (Figs S.46–S.48) (see also LOWER LIMB, PAIN IN, p. 375, Figs L.53–55; BACK, PAIN IN, ‘Spinal tumours’, p. 53).

**Box S.9 Classification of structural scoliosis**

<table>
<thead>
<tr>
<th>Category</th>
<th>Subcategories</th>
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<tbody>
<tr>
<td>Congenital</td>
<td>A failure of Formation, A failure of Segmentation, A combination of both</td>
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<tr>
<td>Idiopathic</td>
<td>Early onset &lt;7 years of age, Late onset &gt;7 years of age</td>
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<tr>
<td>Neuromuscular</td>
<td>Neuropathic</td>
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<td></td>
<td>Upper Motor Neuron:</td>
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<td></td>
<td>– Cerebral Palsy</td>
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<td></td>
<td>– Spinocerebellar degeneration</td>
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<td></td>
<td>– Syringomyelia</td>
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<td></td>
<td>– Spinal cord tumour</td>
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<td>– Spinal cord trauma</td>
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<td>Lower Motor Neuron:</td>
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<td>– Poliomyelitis</td>
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<td>– Traumatic</td>
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<td>– Spinal Muscular Atrophy</td>
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<td>– Myelomeningocele</td>
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<td></td>
<td>– Dysautonomia (Riley-Day syndrome)</td>
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<td>Myopathic</td>
<td>– Arthrogryposis</td>
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<td></td>
<td>– Muscular Dystrophy</td>
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<td>– Duchenne’s</td>
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<td>– Limb girdle</td>
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<td>– Fascioscapulohumeral</td>
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<td>Neuropathic</td>
<td>Fibre-type disproportion</td>
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<td>Congenital hypotonia</td>
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<td>Myotonic dystrophica</td>
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<tr>
<td>Neurofibromatosis</td>
<td>Mesenchymal Disorders</td>
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<td>Rheumatoid Arthritis</td>
<td>Osteochondrodystrophies</td>
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<tr>
<td>Trauma</td>
<td>Sepsis</td>
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<tr>
<td>Metabolic</td>
<td>Tumours</td>
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<td>e.g., osteoid osteoma</td>
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**Figure S.42** Note loss of the lumbar lordosis but absence of vertebral body collapse (a), despite the extent of L4 vertebral body tuberculosis infection and the presence of large psoas abscesses (arrowed) shown by the magnetic resonance imaging scan (b).

**Figure S.43** Adolescent kyphoscoliosis. Thoracic curve convex to the right.
Infantile scoliosis develops before the age of 3 years but unlike the congenital variety (Fig. S.49), there are no vertebral bony abnormalities. There are two types of curve, those that spontaneously resolve and those that progress. Careful follow-up and analysis of the spinal X-rays is required to determine into which group the patient will fall. There is an association with plagiocephaly and sometimes a postural adduction contracture of one hip.
Torticollis (wry neck)
The congenital variety is due to a unilateral contracture of the sternocleidomastoid muscle, and it produces a deformity in which the head is tilted to the ipsilateral side while the chin is rotated to the contralateral side. A rare cause of a torticollis is the Klippel–Feil syndrome, which is a congenital fusion of two or more vertebrae in the cervical region. There may be other associated abnormalities such as a congenital high scapula, or Sprengel’s deformity (see UPPER LIMB, PAIN IN, p. 703, Fig. U.3).

Spondylolisthesis
This is a forward shift of one vertebral body on another. The levels are nearly always L4 on L5, or L5 on the sacrum. The whole lumbar spine therefore slips forward in relation to the sacrum and the pelvis. As a consequence, the lumbar spine appears shortened, transverse skin creases are present, and the sacrum seems to extend upwards towards the waist. A ‘step’ may be present; this is best felt by running a finger down the lumbar spine. There is often associated hamstring tightness.

Spondylolisthesis is subdivided into: ‘dysplastic’, in which the slip is secondary to congenital changes in the upper part of the sacrum and vertebral arch of L5; ‘isthmic’, which accounts for the majority of the cases and is caused by a stress fracture through, or elongation of, the pars interarticularis of the vertebral arch (Fig. S.50); ‘traumatic’, due to an acute fracture; and ‘degenerative’, which occurs as a result of disc degeneration and facet joint osteoarthritis. The term ‘spondylolysis’ refers to a defect of the pars interarticularis but the vertebral bodies remain aligned (i.e. there is no forward slip).
Tenderness of the spine is usually due to local disease of, or injury to, the tissues at the site of tenderness. Such tenderness is always deep, but it may also be associated with cutaneous hyperalgesia. In a second and less important group, the tenderness is partly or entirely cutaneous and is a referred phenomenon found in visceral disease. In testing for spinal tenderness, it is therefore desirable to differentiate between cutaneous and deep tenderness and, in the case of the latter, between tenderness elicited by pressing upon the spinous processes and tenderness in the adjacent muscles. This is because spinal disease is usually accompanied by local muscular spasm, and the muscles thus affected become tender, although they are not themselves the site of the disease. Failure to allow for this fact is the usual explanation of the mistakes – sometimes serious – that are often made in attributing muscle tenderness to a strain or to a rheumatic condition, when in reality it is due to local spasm in response to disease of the vertebrae, the intervertebral discs, or the spinal cord and its membranes.

The chief conditions in which spinal tenderness occurs are summarized in Box S.10.

The investigation of spinal tenderness requires an exhaustive case history, a careful examination and certain special investigations. The history is of particular importance, because not only will it disclose the duration, site and severity of the spinal symptoms, but it will also indicate whether the spinal cord or nerve roots are involved (root pain, girdle sensations, paraesthesiae in the limb, muscular weakness or stiffness, or sphincter disturbances). A systematic interrogation as to general health, previous diseases, and symptoms referable to the other systems of the body may bring out facts relevant to the spinal condition. There is no laboratory procedure that can provide this information, and a further advantage of the historical approach is that it provides a guide to the patient’s mental and emotional condition, which is invaluable in assessing the reality and severity of the spinal symptoms.

The second step, a physical examination, must cover the whole body in a search for factors that may throw light on the spinal tenderness. Reference has already been made to the need for care in determining that the tenderness is really in the spine itself, and not in the adjacent muscles or the overlying skin. (A useful method for examining the vertebrae is to gently ‘spring’ each spinous process, starting proximally and moving distally. To do this, lie the patient down prone, and using the hypothenar eminence of one’s hand, apply a sudden but relatively gentle pressure to each spinous process in turn.) Acute tenderness of organic origin is always associated with limitation of movement in one or more directions. Beware of the spine that has loss of movement in all directions. Assume that such a spine has an infective, inflammatory or neoplastic problem until proven otherwise. The characteristic of mechanical derangement in the spine is that movement in at least one direction remains almost full. For example,
a patient presenting with an acute lumbar disc associated with a lumbar tilt/scoliosis will retain a good range of movement on lateral flexion towards the side of the tilt. The examination of sensation, power and reflexes below the level of tenderness is important as significant neurological abnormalities may be found in the absence of any subjective symptoms. Attention must be paid to the chest, cardiovascular system, abdomen and prostate. The long bones should receive attention, and the skull must not be forgotten, because in carcinomatosis, painless secondary deposits may be found in the latter.

Of the special investigations, X-ray examination of the spine takes the first place. The earliest X-ray changes of pyogenic infection are often fluffiness along the lower border of the body (i.e. at the inferior endplate). The characteristic of established infection is involvement of two bodies and the intervening disc (Figs S.51–S.53). Anatomically, the infection is reflecting the original segmentation, which is half a vertebral body, the disc and half a vertebral body. Metastatic deposits classically show as loss of a pedicle on the anteroposterior film of the spine, and this is easy to miss on a poor-quality X-ray. Lesions in the sacrum are notoriously difficult to appreciate on plain films. Evidence of local disease may be long delayed, and it is dangerous to assume

Figure S.51 *Staphylococcus aureus* infection. This began either as a discitis (left arrow) or an osteomyelitis (right arrow) of the distal aspect of the body of L3.

Figure S.52 Same patient as Fig. S.51. Aspiration under radiological guidance to establish the bacterial diagnosis. Note: a wide variety of organisms can be found in immunocompromised/AIDS patients and drug addicts.

Figure S.53 Same patient as Fig. S.52. Magnetic resonance imaging reveals that the disease is more extensive than was suggested on the plain X-rays. Both bodies, L3 and L4, are involved – ‘two bodies and the intervening disc’ – which is the classical pattern of spinal infection.
that a negative finding is conclusive. CT scanning provides very accurate delineation of the spinal anatomy, especially the bony anatomy. In general, the soft-tissue anatomy is best shown on a magnetic resonance imaging scan.

Check the erythrocyte sedimentation rate (ESR) and the C-reactive protein (CRP) level as well as the haemoglobin, white blood cell count and electrolytes. A raised ESR and CRP requires further thought before attributing backache to a mechanical cause. A rise in the serum acid phosphatase or prostate-specific antigen will suggest the presence of a secondary growth from the prostate, whereas a high alkaline phosphatase is found in Paget's disease. A raised serum calcium level may suggest hyperparathyroidism.

**TENDERNESS IN THE SPINE DUE TO DISEASE IN OTHER PARTS OF THE BODY**

Superficial tenderness over the spine is a common association of visceral disease, and the tenderness is situated over the portion of the spine corresponding to the segmental innervation of the affected viscus. The tenderness is not associated with local rigidity, and there is invariably well-marked evidence of the visceral disease, so that such tenderness should not be mistaken for a manifestation of spinal disease (Figs S.54 and S.55). However, it is easy to be confused, particularly in the case of the elderly, the majority of whom will of course have abnormal findings on plain spinal X-rays. Thus, for example, attributing chest pain to a mechanical problem in the thoracic spine in a 70-year-old is a diagnosis that should only be made by exclusion.

Suspected intercostal nerve root entrapments can be confirmed by use of local anaesthetic. In the abdomen, these nerve roots are classically ‘trapped’ at the edge of the lateral border of the rectus abdominis. A useful physical sign is that the tenderness, which is often diffuse, will localize to this site if the abdominal muscles are contracted by asking the patient to raise the head and torso slightly off the bed while lying supine. In right iliac fossa tenderness, these nerve root entrapments are a differential diagnosis for appendicitis, especially in patients in middle age.

**HYSTERIA, MALINGERING, COMPENSATION NEUROSI**

Psychogenic factors are important to assess, especially in patients with chronic backache (see BACK, PAIN IN, p. 45). Excessive spinal tenderness is one of the ‘inappropriate signs’. Characteristically, even the lightest touch to the skin of the back causes ‘severe pain’. Frequently patients ‘wobble’ and may even lose their balance. By contrast, it is uncommon to find marked tenderness in most cases with serious underlying disease. For example, in spinal infection or ankylosing spondylitis, one may need to ‘spring the spine’ to elicit any tenderness (see above).

**Figure S.54** Spinal metastasis. This patient presented with persistent backache and weight loss. Tenderness was localized over the spinous process of T8. There was a loss of the right pedicle on an anteroposterior X-ray (arrowed).

**Figure S.55** Same patient as Fig. S.54. Magnetic resonance imaging scan showing involvement of the body with early collapse posteriorly onto the cord.
The causes of splenomegaly are numerous, and the most likely diagnosis varies considerably with the age, geographical location and social habits of the patient (Box S.11).

Patients with splenomegaly may present with the symptoms and signs of pancytopenia due to hypersplenism, which may occur with only a modest enlargement of the organ. There is usually an approximately parallel reduction in the number of erythrocytes, leucocytes and platelets, although on occasions there may be a more marked reduction of only one cell line. In the presence of such haematological abnormalities, it is important to demonstrate normal or hyperplastic bone marrow morphology. In addition to the cellular elements being pooled and preferentially destroyed in the spleen, the cytopenias are often exacerbated by a consistent increase in plasma volume.

Many diseases may result in enlargement of the spleen by several different mechanisms. For example, in schistosomiasis, the spleen may be enlarged because of chronic infection as well as portal hypertension secondary to portal fibrosis. A grossly enlarged spleen has a vastly increased blood supply, and portal hypertension may result as a consequence of the splenomegaly. An example is myelofibrosis, which is associated with gross splenomegaly, and at times portal hypertension with ascites. When this is observed, the prognosis is usually poor and splenectomy should be considered.

A careful history and examination will often suggest a diagnosis. A full blood count with differential white-cell count and peripheral blood smear is probably the most important initial investigation. This together with liver function tests and an infectious screen will often provide a short differential diagnosis. An abdominal ultrasound examination will confirm the presence of a mass and also provide accurate dimensions; it is usually possible to distinguish between splenic and the other causes of a left upper quadrant mass (e.g. an enlarged kidney). Furthermore, it may be valuable both for detecting space-occupying lesions (e.g. cysts or abscesses), confirming portal hypertension (Doppler scan of portal vein) and detecting segmental portal hypertension (e.g. due to chronic pancreatitis). An abdominal CT scan with intravenous contrast will provide additional information by delineating the size and consistency of the spleen, and is the investigation of choice if a lymphoma is suspected – retroperitoneal and mesenteric lymph nodes are easily seen.

If the diagnosis is still in doubt, it is usually necessary to undertake fairly wide-ranging investigations. A bone marrow aspirate with biopsy and culture, and a lymph-node or liver biopsy may be needed. A diagnostic splenectomy is occasionally needed but this must be balanced against the short-term operative risks (bleeding, damage to adjacent structures, infections) and long-term risks (infections with encapsulated organisms, thromboembolism, malignancies). Direct splenic puncture appears to be generally safe and permits tissue sampling for infective causes of splenomegaly. It also enables a contrast splenic portogram to be performed if needed.
cough and expectoration; the posture that they adopt may have noticed that certain postures lead to with pleura bronchial fistula. Inquiry can be made as bronchiectasis, lung abscess or rarely an empyema be that it is arising from a localized abnormality, such as eosinophil polymorphs, associated with allergic reactions rather than infection. In acute pulmonary oedema, the material expectorated is derived largely from the oedema fluid transuded into the alveoli; it is thin and frothy, and may be pink from uniform blood-staining. In coal-miners with complicated pneumoconiosis, the sputum contains coal dust. When sputum is profuse, regular and purulent, it can be that it is arising from a localized abnormality, such as bronchiectasis, lung abscess or rarely an empyema with pleura bronchial fistula. Inquiry can be made about the effect of posture upon it; sometimes the patient may have noticed that certain postures lead to cough and expectoration; the posture that they adopt for sleeping may be significant since it may have been chosen because it does not lead to coughing. The sudden production of a large volume of sputum suggests the evacuation of a localized collection of liquid into a bronchus from a pleural empyema, a cyst (infected or otherwise), a lung abscess or a mediastinal, subphrenic or intrahepatic abscess. An episode of this sort may be followed by persistent expectoration or may cease temporarily when the bronchial communication becomes occluded, and recur later. Rupture of a hydatid cyst in the lung may result in the sudden expectoration of a large amount of thin watery material, which may be accompanied or followed by an anaphylactic reaction. Sputum may be odourless, and this gives no indication of the likely pathogens; but if it has a certain type of offensive smell, it is likely that there is an infection with anaerobic organisms, as may occur in some types of lung abscess, empyema with pleurobronchial fistula and severely infected bronchiectasis. The pus in acute specific lung abscesses due to *Staphylococcus aureus* or *Klebsiella pneumoniae*, and in empyemas due to these organisms, to pneumococcus or to *Streptococcus pyogenes*, is not malodorous. In addition to the yellow of pus or eosinophil pseudopus, other colours may be observed in the sputum. A uniformly purulent sputum may be green rather than yellow, either because of degeneration of leukocytes in specimens that have been left standing, or because of infection by *Pseudomonas aeruginosa*, which can be seen as ‘pea green’. The ulceration of an amoebic liver abscess into the lung gives rise to the expectoration of reddish-brown so-called ‘anchovy sauce’ pus. Sputum may be stained with fresh or altered blood, or mixed with larger quantities of blood; this is considered under haemoptysis (see p. 247). The sputum of those exposed to dust will contain the dust that has settled on the bronchi, often aggregated by ciliary streaming to give a mottled appearance. In coal-miners with complicated pneumoconiosis, the confluent collagenous masses incorporating coal dust that constitute progressive massive fibrosis sometimes liquefy centrally; when this occurs, the liquid black contents are expectorated, resulting in an episode of expectoration of inky black material, or ‘melanoptysis’. Formed elements may be visible in sputum. Although careful search by floating the sputum in water may reveal fragments that are evidently casts of small parts of the peripheral bronchial tree in patients with asthma or with diffuse bronchitis, large casts with multiple branching are rarely seen. They occur in the very

Sputum

Alex West

Normally about 100 ml of bronchial secretion is removed daily by ciliary action from the airways of the lung through the larynx, and disposed of into the alimentary tract by unconscious acts of swallowing. Sputum consists of: (i) bronchial secretions in excess of the amount that can be disposed of in this way; (ii) pathological secretions, exudates and pus from abnormal bronchi, bronchioles and alveoli, or from abscesses, cavities or cysts in the lung; or rarely (iii) material derived from morbid processes in pleura, lymph nodes, mediastinum, oesophagus, subphrenic space and liver that have ulcerated into the lung. It may be mixed with saliva and secretions from the upper respiratory tract but should be distinguished from these. It should be remembered that the commonest cause of purulent sputum is a simple lower respiratory tract infection or bronchitis. The patient's account of its mode of production and the quantity and quality of the sputum, and the physician's observations, especially the naked-eye appearances, often give information at least as important as that derived from laboratory procedures. Although the production of sputum is usually associated with cough, some patients with chronic bronchitis deny a cough, regarding the expectoration of mucus or even mucopus in the mornings as so ‘normal’ that they refer to it as ‘clearing their throat’. Other patients, who evidently raise excess secretions from their lower respiratory tracts by cough, deny producing sputum because they habitually swallow the material expectorated.

Sputum arising from the bronchi may be mucoid, mucopurulent or frankly purulent. It is important to remember that, in asthmatic patients, yellow sputum does not necessarily indicate the presence of pus (i.e. neutrophil polymorphs); it may be due to eosinophil polymorphs, associated with allergic reactions rather than infection. In acute pulmonary oedema, the material expectorated is derived largely from the oedema fluid transuded into the alveoli; it is thin and frothy, and may be pink from uniform blood-staining. When sputum is profuse, regular and purulent, it can be that it is arising from a localized abnormality, such as bronchiectasis, lung abscess or rarely an empyema with pleura bronchial fistula. Inquiry can be made about the effect of posture upon it; sometimes the patient may have noticed that certain postures lead to cough and expectoration; the posture that they adopt...
rare plastic or fibrinous bronchitis; the patient, who is often asthmatic, suffers recurrent febrile illnesses with collapse–consolidation of a lobe or lobes of the lung, re-expanding after expectoration of the cast. Much more frequent is allergic bronchopulmonary aspergillosis, in which the sputum may contain ‘plugs’, generally about 4–5 mm in diameter and 15–20 mm long. Usually, they are roughly spindle-shaped, without the multiple branching of bronchial casts, although there is occasionally a single bifurcation at one end. They consist mainly of tough mucus, containing many eosinophils, and with a little Aspergillus mycelium in the centre, usually demonstrable only by special staining. This disease occurs in extrinsic atopic asthmatics, and the sputum may also have the microscopic features seen in asthma (see below). A careful search in the sputum of asthmatic patients suspected of allergic aspergillosis may be required to demonstrate the ‘plugs’, but patients will on inquiry often be found to have noticed the presence from time to time of a tough fragment in the sputum. Very rarely, a patient with a bronchial carcinoma coughs out a gross fragment of the tumour. Another rare event is the expectoration of a fragment of calcified caseous material from an old tuberculous focus, either in lung or in a bronchopulmonary lymph node. If a previous chest radiograph is available, it is sometimes possible to see that one of the calcified foci evident in it has disappeared in a subsequent film.

The principal laboratory examinations to which sputum should be submitted are microscopy and bacteriological culture.

The presence of pus can be microscopically confirmed by the finding of large numbers of neutrophil polymorphs. As already noted, it is important to distinguish the eosinophil pseudopus which appears in the sputum of some asthmatics from true pus. Additionally, in the mucoid sputum of asthmatics, many eosinophils may be present. This finding is of particular importance in the differential diagnosis between late-onset intrinsic asthma and chronic bronchitis. The sputum of asthmatic patients may also contain Curschmann’s spirals and Charcot–Leyden crystals. Curschmann’s spirals consist of whitish, twisted threads of mucus, often including eosinophils; Charcot–Leyden crystals are colourless, elongated octahedrons that appear to be associated with eosinophils. Occasionally, small clumps of desquamated bronchiolar epithelial cells, the so-called ‘Creola bodies’, may be seen in the sputum of asthmatic patients, especially after a severe attack or during a prolonged attack.

Examination of the sputum for cancer cells can sometimes be a helpful investigation in the diagnosis of bronchial carcinoma, but it is now not widely used as bronchoscopy has a far higher diagnostic rate. Clinicians should be aware of some possibly confusing factors. In asthmatic patients, a report that clumps of adenocarcinoma cells have been seen should be interpreted in the knowledge that Creola bodies, mentioned above, may mimic such cells very closely; and in patients with chronic tuberculous or other cavities in the lung, which may be lined with metaplastic squamous cells, these cells may be desquamated and prove difficult to distinguish with certainty from squamous carcinoma cells.

After haemoptysis from any cause and in the presence of pulmonary congestion associated with heart disease, iron-containing macrophages, or siderocytes, may be seen in the sputum. They also appear in the sputum in idiopathic pulmonary haemosiderosis but are not of specific significance in this disease.

Persons who have been exposed to asbestos dust produce ‘asbestos bodies’ in their sputum. These consist of very thin, needle-like fibres of asbestos surrounded by a clear brownish coating of proteinaceous material containing iron, often arranged in an irregular beaded distribution, or with a terminal bead or beads causing the whole to resemble a drumstick or dumb-bell. The presence of these bodies indicates only exposure to asbestos and is not necessarily associated with pulmonary asbestosis. Apart from this, microscopy of the sputum gives no specific information in pneumoconioses. Similarly, although oil-containing macrophages may be found in the sputum of patients with exogenous oil inhalation pneumonia, they may also be found in users of oily nasal drops – but without pathological consequences in the lungs.

In pulmonary alveolar proteinosis, microscopy of the sputum shows amorphous eosinophilic periodic acid–Schiff-positive material, while electron microscopy shows the presence of lamellar bodies, presumably derived from type II pneumocytes, which may be diagnostic.

Microscopy of suitably stained sputum smears is an essential part of the examination of the sputum for mycobacteria. It has been estimated that sputum specimens must contain as many as 100,000 bacilli per ml if acid-fast bacilli (AFBs) are to be reliably demonstrated in them by microscopy after Ziehl–Neelsen staining. Examination by fluorescence microscopy after suitable staining has a somewhat higher sensitivity. AFBs seen on staining means that...
mycobacteria are present, but culture is required to determine if it is *M. tuberculosis* or a non-tuberculous mycobacterium (NTM). Certain NTMs can be pathogenic in their own right but there are a large number of NTMs which may be considered ‘environmental’ and non-pathogenic to humans. Appropriate methods of culture demonstrate mycobacteria in specimens containing far fewer organisms, but there is a delay of several weeks before the result can be available. For this reason, persistent attempts should be made to find acid-fast bacilli by microscopy in any patient who is acutely ill with an inflammatory process in the lung that might be tuberculous. In this interpretation of negative findings, it is important to remember that failure to find acid-fast bacilli in a scanty mucoid sputum in a patient with acute pneumatic changes without cavitation militates very little against a diagnosis of tuberculosis, whereas in a patient with a cavitated inflammatory process and frankly purulent sputum, repeated negative findings might be more significant.

Microscopy of Gram-stained smears of sputum is of value in acute pneumonias – for example, a preponderance of Gram-positive diplococci suggests a *pneumococcal infection*, or of clumps of Gram-positive cocci a *staphylococcal infection* – but, in bacterial infections, culture is generally required both to identify organisms and to provide information about their sensitivity to antibiotics. Sputum is usually cultured only aerobically. Culture of expectorated sputum anaerobically is useless because the sputum is inevitably contaminated by oropharyngeal organisms, which include many anaerobic species. If infection with anaerobic organisms is suspected, specimens must be obtained from the lower respiratory tract either by transtracheal aspiration or at fibreoptic bronchoscopy using a sheathed brush to obtain the specimens.

In patients suspected of *pneumocystis pneumonia*, specimens of alveolar secretion may be obtained by alveolar lavage and examined by immunofluorescent staining techniques for *Pneumocystis jiroveci* (formerly *Pneumocystis carinii*). It should be remembered that, in the sputum of patients receiving broad-spectrum antibiotics, organisms other than the original pathogens often become predominant. These may be organisms that rarely become pathogenic, such as *Proteus* and coliform organisms, as well as some that can assume independent pathogenicity, such as *Pseudomonas pyocyanea* and *Candida*. Even with the latter, it is often difficult to be certain whether or not they are truly pathogenic in an individual case of chronic mucopurulent bronchitis.

Sporing organisms, such as *Aspergillus* species, the spores of which are frequently present in the air, appear as contaminants in a proportion of all sputum cultures. For this reason, the finding of an *Aspergillus* species in the sputum is of no significance unless supported by clinical and immunological evidence. When infection with other pathogenic fungi is suspected, special culture media are required; such cultures can, if positive, be highly infectious and a laboratory hazard.

**SQUINT (STRABISMUS)**

Reginald Daniel

Squints may be classified according to their direction, into convergent (Fig. S.56), divergent or vertical; and according to their cause into paralytic and non-paralytic (concomitant). The differential diagnosis between a paralytic and a non-paralytic squint is, as a rule, straightforward. In a paralytic squint, the degree of deviation of the two eyes varies, as the further the eyes are moved in the direction of the action of the paralysed muscle, the greater will be the angle of squint. In a concomitant squint, the eyes always bear the same relative position to each other in whichever direction they are turned. Concomitant squint is characteristically a disorder of childhood, while paralytic squint more frequently occurs later in life.

The diagnosis of the cause of a paralytic squint is discussed under DIPLOPIA (p. 138). Concomitant squints are usually the sequel to disharmony of the accommodation–convergence synkinesis (as with high hypermetropia), poor development in the power of fusion, or anatomical fasciomuscular abnormalities of the eyes – all of which may have a congenital basis. They may be aggravated by any sensory or central impediment.
to the acquisition of the binocular fixation reflex, for example poor vision from a congenital cataract, or poor coordination from learning difficulties.

**STATURE, SHORT**

Paul Carroll

Patients can be described as suffering from ‘short stature’ when they are shown to be below the 3rd centile of a normal population of their sex and age. Charts have been devised by Tanner and Whitehouse for a normal British population, but these are not necessarily applicable to people of a different ethnic origin. Most short British children are so because they have small parents, are socially deprived or are suffering from delayed puberty, whereas in developing countries of the world, common causes of short stature are malnutrition and chronic debilitating disease in childhood. Rarer causes include skeletal deformities and hormonal deficiencies. The causes of short stature are listed in Box S.12.

**NORMAL GENETIC SHORT STATURE (NORMAL VARIANT SHORT STATURE, NVSS)**

Most children who present with short stature are short because they have small parents. This can be readily ascertained by plotting the child’s height on a growth chart for the relevant population and comparing it with the parents’ heights.

The short child whose centile falls within his or her parents’ normal range should be seen in 6 months’ time, to make sure that the growth velocity is normal.

**GROWTH DELAY (NORMAL VARIANT CONSTITUTIONAL DELAY, NVCD)**

Growth delay, which is often associated with delayed puberty, is a common cause of short stature. There is often a family history of both. The bone age, assessed from a radiograph of the wrist and hand, will usually be delayed by 3 years or more. Ultimately, normal growth and sexual development is the rule.

**CHROMOSOMAL ABNORMALITIES**

The most common chromosomal abnormalities are Turner’s syndrome and Down’s syndrome.

**Turner’s syndrome**

Turner’s syndrome occurs in girls and is characterized by gonadal dysgenesis, a lack of sexual development and short stature. It occurs with a prevalence of 1 per 3000 female births. The karyotype is 45,XO, and the buccal smear will show no Barr bodies. The average height reached is 140 cm, and rarely is a height above 152 cm achieved. However, about one-quarter of all

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**Box S.12 Causes of short stature**

<table>
<thead>
<tr>
<th>Normal genetic short stature</th>
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<tbody>
<tr>
<td>Growth delay</td>
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<tr>
<td>Chromosomal abnormalities</td>
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<tr>
<td>Gonadal dysgenesis</td>
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<tr>
<td>- Turner’s syndrome</td>
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<tr>
<td>- Noonan’s syndrome</td>
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<tr>
<td>Autosomal anomalies</td>
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<tr>
<td>- Trisomy 21 (Down’s syndrome)</td>
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<tr>
<td>- Trisomy 18 (Edward’s syndrome)</td>
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<td>- Trisomy 13 (Patau’s syndrome)</td>
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<tr>
<td>Disease in childhood</td>
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<tr>
<td>- Calorie deficiency (marasmus)</td>
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<tr>
<td>- Protein malnutrition (kwashiorkor)</td>
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<td>- Vitamin D deficiency (rickets)</td>
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<tr>
<td>- Tuberculosis</td>
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<tr>
<td>- Bronchiectasis</td>
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<tr>
<td>- Cystic fibrosis</td>
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<tr>
<td>- Chronic asthma (particularly when treated with steroids)</td>
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<tr>
<td>- Gluten enteropathy (coeliac disease)</td>
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<tr>
<td>- Other malabsorption syndromes (e.g. Crohn’s and ulcerative colitis)</td>
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<tr>
<td>- Hookworm infection</td>
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<tr>
<td>- Malaria</td>
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<tr>
<td>- Cyanotic congenital heart disease</td>
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<td>- Chronic renal disease</td>
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<tr>
<td>- Congenital syphilis</td>
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<td>- Glycogen storage disease</td>
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<tr>
<td>- Thalassaemia major</td>
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<tr>
<td>Psychological causes</td>
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<tr>
<td>Skeletal abnormalities</td>
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<tr>
<td>Congenital</td>
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<tr>
<td>- Achondroplasia, hypochondroplasia</td>
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<td>- Morquio’s disease (chondro-osteodystrophy)</td>
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<td>- Dysostosis multiplex (gargoylism or Hurler’s syndrome)</td>
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<tr>
<td>- Chondrodystrophy calcificans congenita (Conradi’s syndrome)</td>
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<tr>
<td>- Epiphysial dysplasia multiplex</td>
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<tr>
<td>- Chondro-ectodermal dysplasia (Ellis–van Creveld syndrome)</td>
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<tr>
<td>- Osteogenesis imperfecta (fragilitas ossium)</td>
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<tr>
<td>- Approximately 50 others</td>
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<tr>
<td>Acquired</td>
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<tr>
<td>- Rickets (see below)</td>
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<tr>
<td>- Tuberculosis and other infections of the spine</td>
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<tr>
<td>- Deformities secondary to neurological and joint diseases (e.g. poliomyelitis, Still’s disease)</td>
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<tr>
<td>Endocrine abnormalities</td>
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<tr>
<td>- Growth hormone deficiency (GHD)</td>
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<tr>
<td>- Familial</td>
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<tr>
<td>- Type IA (complete, autosomal recessive)</td>
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<tr>
<td>- Type IB (incomplete, autosomal recessive)</td>
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<td>- Type II (autosomal dominant)</td>
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<tr>
<td>- Type III (X-linked GHD)</td>
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(Continued)
patients with Turner’s syndrome have mosaicism, such as XO,XX or XO,XXX, and they may grow to a greater height. These patients may show one or, in the latter case, even two Barr bodies on buccal smear. It is wise in such circumstances to perform a full chromosome analysis.

Because the ovaries are represented only by a fibrous streak, no oestrogens are secreted, so both the internal and external genitalia remain infantile. No breast development occurs, and there is primary amenorrhoea. In addition, there are various physical abnormalities that are not hormonally mediated. The short stature is in contrast to most other types of primary hypogonadism, in which tall stature due to a lack of epiphysial fusion is a feature. There may be webbing of the neck and a low hairline at the back of the neck. There is an increased carrying angle at the elbows (cubitus valgus), shortened fourth and/or fifth metatarsals and metacarpals, Madelung’s deformity of the wrist (abnormal carpal angle), a shield-like chest and abnormal fingerprints (dermatoglyphics). Pigmented naevi are often present on the skin, and intestinal telangiectasia sometimes occurs. Congenital cardiovascular anomalies exist in about one-fifth of cases; these consist of coarctation of the aorta, aortic stenosis and both atrial and ventricular septal defects. Renal abnormalities such as horseshoe kidney or double ureter may occur. Hypertension is not uncommon. Subcutaneous oedema may be present in infancy. Minor abnormalities of the facies frequently draw attention to the diagnosis. The features include a flattened bridge to the nose, wide separation of the eyes (hypertelorism) and epicanthic folds. Osteoporosis may develop late in life due to the lack of oestrogens. Autoimmune thyroid disease is more common in patients with Turner’s syndrome than it is among the general female population.
Noonan’s syndrome
This syndrome may occur in boys as well as girls and is somewhat like Turner’s syndrome. The prevalence is 1 in 8000 births. In both boys and girls, the karyotype is normal. Short stature is perhaps not so marked as in Turner’s syndrome, but there may be webbing of the neck with a low hairline and a shield-like chest with widely spaced nipples. Pectus excavatum is present in 50 per cent of cases. The face shows some typical features: a slant of the eyes, which are widely spaced, may have epicanthic folds and show ptosis; the face may be triangular in shape, the ears low-set and the brow prominent. A high-arched palate is not infrequently present. The cardiovascular abnormalities differ in frequency from those of Turner’s syndrome. Although coarctation of the aorta and aortic stenosis have been described, the characteristic cardiac diseases have been pulmonary stenosis (in approximately 50 per cent of cases) and atrial septal defect. Sometimes, both conditions occur together. Ventricular septal defect and persistent ductus arteriosus have also been associated with Noonan’s syndrome. Cubitus valgus is common, and mental retardation may be a feature. In males, undescended testes are common, and androgen deficiency may manifest itself at puberty. Girls usually show delayed puberty, but eventually normal ovarian function develops.

Down’s syndrome (trisomy 21)
This is the most common autosomal abnormality leading to short stature. There is disordered growth of the skull and long bones. The mouth and ears are small, the tongue is large and tends to protrude, and there are epicanthic folds on the eyes. There is invariably mental retardation. Autoimmune thyroiditis is much more common than in the general population.

Trisomy 18
Trisomy 18, also known as Edward’s syndrome, is the second most common autosomal condition, and is characterized by a small face and cardiac malformations that nearly always lead to an early death. It occurs with a prevalence of 1 per 8000 births.

Trisomy 13
Trisomy 13, also known as Patau’s syndrome, is much more rare, and occurs only once in 20000 live births. Death tends to occur in the first year of life.

MALNUTRITION AND CHRONIC DEBILITATING DISEASE IN CHILDHOOD
General calorie deficiency (marasmus) and protein deficiency (kwashiorkor) are common causes of short stature in underdeveloped parts of the world. Rickets, which is now unusual among the indigenous population of Britain, but relatively frequent among Asian children resident in this country, leads to retarded bone age and short stature. Self-inflicted malnutrition due to anorexia nervosa and bulimia are also common in the developed world. Growth failure in these circumstances is characterized by a normal or elevated growth hormone (GH) with decreased somatomedin levels.

The existence of chronic debilitating disease is readily apparent on taking the patient’s history or performing the physical examination. Tuberculosis and bronchiectasis are not nearly as common as they used to be in developed countries. Cystic fibrosis occurs in approximately 1 in 1700 live births among Caucasians, but it is much rarer in other ethnic groups. Recurrent pulmonary infections are usually the predominant feature. Chronic asthma, particularly when treated with steroids, leads to stunting, probably because excess glucocorticoids block the action of growth factors at the cellular level.

Malabsorption of food may lead to short stature. Gluten enteropathy (coeliac disease) is the most common such condition, occurring in approximately 1 in 2000 children. Although characteristic symptoms of chronic diarrhoea, pale bulky stools, anorexia and cramping abdominal pain may be present, they are by no means invariably. Sometimes, the only clue in a short child is a rather protuberant abdomen and scanty subcutaneous fat. If other causes of short stature have been excluded, a small-bowel biopsy to show flattened villi is a useful investigation, as, once the diagnosis has been confirmed and a gluten-free diet instituted, normal growth should occur. Less common causes of malabsorption include cystic fibrosis (see above), lactose intolerance, Crohn’s disease and infection with Giardia lamblia.

Children who either live in the tropics or have recently returned from there may have hookworm infestation or malaria. Thalassaemia major may be associated with short stature and delayed puberty.

Cyanotic congenital heart disease and chronic renal failure are usually obvious clinically as causes of growth failure. Less commonly seen are congenital syphilis and glycogen storage disease.

PSYCHOLOGICAL CAUSES
An adverse psychological environment, with emotional deprivation and child abuse, is a very important cause of failure of growth. This has been shown to be due to suppression of GH secretion by the pituitary, but nutritional factors undoubtedly play a role. It is reversible when the child is removed to happier and more secure surroundings.
SKELETAL DISORDERS

Congenital skeletal disorders

These are rare, and many of them are inherited. The most common is achondroplasia (chondrodys trophy). The condition, which is autosomal dominantly inherited, is caused by a disturbance of endochondral ossification in which the growth cartilages are invaded by connective tissue. The trunk is of normal length, but the limbs are shortened. The fingers are short and are of equal length. The head is large, the forehead being particularly prominent, but the nose is small and the bridge is flattened. There is often a marked lumbar lordosis, and the buttocks are prominent. Delayed puberty is not a feature.

Hypochondroplasia is similar to achondroplasia, but the face is usually normal. Again, the inheritance is autosomal dominant.

Morquio’s disease (chondro-osteodystrophy) is a rare type of skeletal deformity affecting both the limbs and the spine, and it is often familial. The epiphyses are deformed or fragmented, and a variety of deformities of the long bones have been described. The glenoid fossae and acetabulae are poorly formed. The intervertebral spaces are widened and, owing to the irregularity, flattening or wedge-shaped deformity of the vertebral bodies occurs, and there is a dorsolumbar kyphosis and shortening of the neck, the head appearing to be pushed down onto the shoulders.

Dysostosis multiplex (gargoylism or Hurler’s syndrome) shows similar bony deformities to the above, although a peculiar sabot-shaped deformity of the second and third lumbar vertebral bodies appears more constantly, giving rise to angular kyphosis. The skull commonly is conical (oxycephalic) or hydrocephalic, and there may be enlargement of the pituitary fossa. The face shows widely spaced eyes (hypertelorism), a prominent forehead and a large tongue and lips. The liver and spleen may be enlarged. Mental retardation, corneal opacities and deafness are the rule. Distortion of the heart valves and thickening of coronary arteries are frequent findings. The condition is inherited as an autosomal recessive trait and is due to the deposition of the mucopolysaccharides heparan sulphate and dermatan sulphate. Infantilism may be associated with the short stature, but this is not always the case. However, most children do not survive beyond the age of about 14 years.

Chondrodys trophya calcificans congenita (Conradi’s syndrome) is a rare autosomal recessive condition in which the epiphysial centres of the long and small bones ossify and fuse during early childhood.

Epiphyseal dysplasia multiplex can be transmitted as an autosomal dominant or as a recessive type of inheritance. It manifests itself in older children (usually under the age of 10 years) as a disorder of gait and short stature. The pathology lies in the epiphyses, which are fragmented.

Chondro-ectodermal dysplasia (Ellis–van Creveld syndrome) causes short stature because of a reduction in length of the extremities. The fingers are also short, and there may be polydactyly. Hypoplasia of the teeth, nails and hair occurs, and there may be both cardiac and renal abnormalities.

Osteogenesis imperfecta (fragilitas ossium) is characterized by extreme brittleness of the bones. The multiple fractures that occur during intrauterine life or in childhood, coupled with the fragility of the spine, result in gross deformity and short stature. The disease is inherited as an autosomal dominant trait, and there may be a family history of otosclerosis. The patient may have slate-blue sclerae. Radiographically, the long bones are seen to be both shorter and more slender than normal, and poorly calcified, with extreme thinning of the cortex. The appearances are quite distinct from those of rickets.

Acquired skeletal disorders

Any disorder leading to damage of the long bones or spine will lead to stunted growth. Rickets (see p. 644) is not uncommon among Asian immigrant communities in Britain, particularly those who are vegetarians. Tuberculosis of the spine (Pott’s disease, spinal caries) is much less common than it used to be. It is recognized by pain and tenderness localized to one or more of the vertebrae, associated with angular kyphosis. Radiographs show destruction and collapse of the affected vertebral bodies, and there may be evidence of a ‘cold abscess’. These features serve to distinguish spinal tuberculosis from congenital abnormality of the vertebrae and kyphosis or scoliosis from other causes (e.g. poliomyelitis), in which weakness of the muscles of the back, perhaps unrecognized, is liable to result in a severe degree of postural deformity.

Still’s disease (chronic juvenile arthritis; see p. 333) may lead to a stunting of growth because of premature fusion of epiphyses in affected joints.

ENDOCRINE DISORDERS

Endocrine abnormalities account for approximately 10 per cent of cases of short stature. It is important to make the diagnosis promptly in children as
appropriate therapy can normalize growth. The most common endocrine causes of short stature are growth hormone deficiency (GHD), which may be partial rather than total, and hypothyroidism. Less common are suprarenal disorders, for example glucocorticoid excess, either iatrogenic or secondary to Cushing’s disease, and primary suprarenal hypercortisolism. The remaining causes listed in Box S.12 are much more rare.

**Growth hormone deficiency** is a mixture of genetic abnormalities. GHD is recognized early in life, because the children tend to be fat with small genitalia, short stature and hypoglycaemia. They have delayed bone age and delayed puberty, but they enter puberty when their bone age is appropriate. There are two variants of type 1 GHD, depending on the degree of genetic mutation. Type 1A GHD is due to gene deletion or point mutations, and there is complete GHD. These patients develop GH antibodies when treated. Type 1B GHD is associated with splice-site mutations in the GH gene. Patients retain the ability to produce small amounts of GH, and they do not develop antibodies when treated with GH. Destructive lesions involving the pituitary and hypothalamus may cause panhypopituitarism. These include tumours, of which the most common in childhood is a craniopharyngioma, but other causes are meningitis (e.g. tuberculous), sarcoidosis, xanthomas (e.g. Hand–Schüller–Christian syndrome or histiocytosis X), trauma (especially basal skull fractures) and following intracranial irradiation and/or chemotherapy. A subset of children with normal GH responses to stimulatory testing may still respond to GH treatment. These children are thought to be suffering from a neurosecretory dysfunction of GH secretion. A more comprehensive system of interpreting GHD may therefore come from the combination of short stature, decreased growth velocity, delayed bone age and low serum insulin-like growth factor (IGF)-I in addition to low levels of IGF binding protein (IGFBP)-3 and another binding protein called the acid-labile subunit. Collectively, the IGFs have been known as the somatomedins, and they play a role in somatic growth on binding of GH to its receptor.

*Fröhlich’s syndrome* is mentioned if only to be dismissed. Fröhlich originally described a patient with obesity, somnolence, retarded skeletal and sexual development, and optic atrophy. The pathology in this case was a suprasellar tumour pressing on the hypothalamus, and clearly pituitary hormone deficiencies secondary to hypothalamic damage were likely to have been present. Unfortunately, the term ‘Fröhlich’s syndrome’ tends to be attached to any short, fat boy with delayed sexual development and small genitalia. It is best forgotten.

Laurence–Moon–Biedl syndrome may sometimes be associated with short stature. The condition is autosomal recessive. Clinical features include obesity, retarded sexual development, mental retardation, retinitis pigmentosa and either polydactyly or syndactylyism.

The rare Laron type of short stature is caused by mutations or deletions of the GH receptor gene and is associated with increased GH levels and decreased levels of IGFs and IGFBP3. It can now be diagnosed by measuring levels of GH-binding protein and can be treated with IGF-I.

The Pygmies of Africa have normal GH and somatomedin levels; an absence of somatomedin receptors has been postulated.

**Hypothyroidism** is a common cause of short stature. Somatomedin levels are below normal. Among babies born in Britain, 1 in 3300 is hypothyroid, usually due to thyroid agenesis. The diagnosis is now made early by neonatal thyroid-stimulating hormone (TSH) screening. However, it must be noted that neonatal hypothyroidism may be secondary to isolated TSH deficiency. This will be missed on TSH screening but will be diagnosed if serum thyroxine is measured. In addition, hypothyroidism may co-exist with GH deficiency and may render false-negative stimulatory tests of GH production. It is important, therefore, to rule out hypothyroidism in the first instance. Dyshormonogenesis, which may be due to one of six defects in the synthesis of thyroid hormones, tends to lead to goitre and hypothyroidism in early life, although those with partial defects may present later. After the age of round about 6 years, autoimmune thyroiditis begins to become an increasingly important cause of hypothyroidism. A goitre is usually palpable, and thyroid antibodies are present in blood. Sometimes at the time of puberty, an ectopic, often sublingual, thyroid may be the cause of hypothyroidism.

Uncontrolled type 1 diabetes mellitus is associated with decreased somatomedin levels and poor growth. Restoration of the growth rate occurs with improved nutrition and diabetes control.

When precocious sexual development occurs, the production of sex hormones initially causes an acceleration of growth, but bone maturation is more rapid than normal, fusion of the epiphyses takes place, and the final stature is shorter than would be expected from parental height (Fig. S.57). Precocious sexual
development is five times more common in girls than in boys, and in the majority of girls the cause is some unknown triggering mechanism in the hypothalamus that initiates a normal, albeit early, physiological puberty. In boys, on the other hand, a pathological cause is found in about half the cases.

It is important to separate true precocity from false precocity. In true precocity, puberty is ‘physiological’ and thus follows the pattern of normal development. While the majority of cases are constitutional, it is important to look for space-occupying lesions of the central nervous system. These include craniopharyngioma, pineal tumour, hamartoma, neurofibroma, astrocytoma and cysts of the third ventricle. Hypothyroidism may cause the premature release of follicle-stimulating hormone (FSH) from the pituitary. Tuberous sclerosis, McCune–Albright syndrome (polysostotic fibrous dysplasia with cutaneous pigmentation), encephalitis and hydrocephalus may all lead to true precocity. On the other hand, ‘false’ precocity is due to the action of sex hormones produced by the gonad or by the suprarenals. The common causes are congenital adrenal hyperplasia due to a 21-hydroxylase defect, and Cushing’s syndrome. Testicular and ovarian tumours are very rare. For further details, see PUBERTY, PRECOCCIOUS (p. 535).

*Pseudohypoparathyroidism* is a disorder in which the renal tubules are resistant to the action of parathyroid hormone. As a result, the production of 1,25-dihydroxy-vitamin D by the kidneys is reduced and, as a consequence, the absorption of calcium from the gut is diminished. The patient is usually short and mentally retarded, has short fourth and fifth metacarpals, and may have cataracts and ectopic subcutaneous calcification with ulceration. Serum calcium levels are low, and parathyroid hormone levels are increased. Some patients have the short stature and skeletal abnormalities but with a normal calcium level. This is called ‘pseudo-pseudohypoparathyroidism’.

*Rickets* may lead to stunting of growth and bowing of the legs. The most common cause is vitamin D deficiency due to lack of ultraviolet light, so that the precursor of the vitamin, 7-dehydrocholesterol, is not converted into vitamin D. The normal British diet contains very little vitamin D. Groups particularly at risk are Indian and Pakistani immigrants, especially if they are vegetarian (Fig. S.58). Small-bowel and biliary diseases also cause malabsorption of vitamin D. Other rarer causes of rickets are congenital renal tubular defects and 25-hydroxyvitamin D-1-alpha-hydroxylase deficiency (vitamin D-resistant rickets).

**Figure S.57** Comparison of short stature due to early closure of epiphyses, and tall stature due to late closure of epiphyses because of hypogonadism, but with normal growth hormone secretion. [Courtesy of the Departments of Medical Illustration, Guy’s and Westminster Hospitals.]

**Figure S.58** Rickets in a vegetarian Indian boy. Note the genuvalgum.
INTRAUTERINE MALDEVELOPMENT

Most small-for-date babies subsequently grow at a normal rate, although they may not achieve a final stature appropriate for their parents' centiles. Some, however, fail to grow normally and may be examples of specific syndromes, of which the most common is the Russell–Silver syndrome. This is characterized by a lack of subcutaneous fat, a triangular face with a large forehead, small jaw, low-set ears and a turned-down mouth. There is incurving and shortening of the fifth fingers.

The fetal alcohol syndrome is being increasingly recognized; this is probably related to the rising consumption of alcohol. In addition to growth retardation, there is mental retardation and a characteristic facial appearance. This consists of a small jaw, short nose, underdeveloped upper lip and reduction in length of the palpebral fissures. More serious conditions are congenital heart defects and flexion contractures.

Progeria (Hutchinson–Gilford syndrome) is characterized by premature ageing, and death usually occurs before the age of 20 years. Appearances are normal at birth, but growth ceases – owing to epiphyseal closure – within a period ranging from a few months to 3 years. There is no subcutaneous fat, the nose is beaky and the ears are prominent. There may be premature baldness, some mental retardation, periarticular fibrosis and arteriosclerosis.

Cornelia de Lange syndrome (Amsterdam dwarfism) is characterized by poor growth, mental deficiency, a small head with a flattened occiput, abnormal lips and mouth, bushy eyebrows that meet in the middle, short nose with anteverted nostrils and abnormalities of the extremities, particularly of the upper limbs. Hirsutism may be a feature.

Patients with Cockayne syndrome have a somewhat similar appearance to that of progeria, except that mental retardation is more marked, and there is retinal degeneration, photosensitive dermatitis, deafness, unsteady gait and tremor.

Prader–Willi syndrome is characterized by poor growth, obesity, hypocondrosism, cryptorchidism, hypotonia and mental retardation. The hands and feet are small in relation to the rest of the body.

DRUGS

The administration of various drugs, particularly in high doses, may lead to stunting of growth in a child. Glucocorticoids, used particularly in asthma and Still’s disease, may cause short stature due to interference with growth at the cellular level. The dosage should therefore be kept to the minimum possible and should preferably be given on alternate days.

Anabolic steroids have sometimes been used to try to increase the height of a short child. The danger is that they accelerate bone age, cause epiphyseal fusion, and may thus actually decrease potential height. The same may happen when androgens or oestrogens are used in boys and girls, respectively, to bring about sexual maturation. Specialist advice should be sought before using any of these drugs in children.

A lack of thyroid hormone (hypothyroidism) causes delayed bone maturation and short stature, but it must be remembered that the condition may be iatrogenic, when too much anti-thyroid drug is given, and that iodides may lead to goitre formation and hypothyroidism in some individuals who are predisposed to autoimmune thyroid disease. Conversely, overtreatment of hypothyroidism in childhood with thyroxine will lead initially to a growth spurt due to stimulation of somatomedin production by the liver but, by accelerating bone maturation, may reduce potential stature in the long term.

Vitamin D preparations, given to children for conditions such as hypoparathyroidism, pseudohypoparathyroidism and rickets, may give rise to hypercalcaemia, which, if chronic, will cause renal failure. If the renal failure is severe enough, stunting of growth will result. Vitamin D treatment should always be monitored with regular measurement of the serum calcium.

STATURE, TALL

Mark Kinirons

A patient is usually defined as exhibiting tall stature if the height is above the 97th centile for the normal population. In most cases, tallness is inherited from tall parents; pathological causes are much less common. However, a child may be below the 97th centile but still be inappropriately tall compared with the parents. If a period of observation shows that the growth velocity is greater than normal, the appropriate investigations should be carried out. The classification of tall stature is listed in Box S.13.

FAMILIAL TALL STATURE OR CONSTITUTIONAL ADVANCED GROWTH

Familial tall stature can be identified by plotting the patient’s height relative to age on a growth-development chart relevant to the child population and ethnicity. Some children who enter puberty relatively early may initially become taller than their contemporaries because of an early growth spurt, but...
thyrotoxicosis

A similar growth acceleration is seen in early, and a normal adult height is not achieved. Unfortunately, epiphyses fuse at an earlier age than usual normalizes the final height. The GH levels may be low, but GH-binding protein and IGF-I levels are normal to high.

Overnutrition leading to obesity in early childhood may lead to accelerated linear growth, but fusion of the epiphyses at an earlier age than usual normalizes the final height. The GH levels may be low, but GH-binding protein and IGF-I levels are normal to high.

ENDOCRINE ABNORMALITIES

The most common endocrine causes of tall stature in childhood are true precocious puberty and false precocious puberty (see p. 535). True precocious puberty is either physiological (which is the most common) or is due to space-occupying or other lesions affecting the hypothalamus. False precocious puberty can be caused by either suprarenal disease (congenital adrenal hyperplasia, e.g. 21-hydroxylase defect, and suprarenocortical tumour) or by very rare hormone-secreting testicular or ovarian tumours. The increased amounts of sex hormones produced in these various conditions lead to an increased maturation of bone, with an acceleration of growth so that the child is inappropriately tall compared with peers of the same age group. Unfortunately, epiphyses fuse early, and a normal adult height is not achieved.

A similar growth acceleration is seen in thyrotoxicosis and in hypothyroid patients who are overtreated with thyroxine. Somatomedin levels are high. Thyrotoxic children are usually round about the 75th centile for height. However, tall stature is rarely a presenting feature in these children, and ultimate stature is not usually abnormal because the epiphyses fuse early. Hypogonadism will cause increased stature with long arms and legs (eunuchoidism), because of delayed fusion of the epiphyses, as long as normal amounts of GH and thyroid hormone are present. Disease of the hypothalamus, pituitary or gonad may cause hypogonadism, leading to tall stature. Solitary deficiency of hypothalamic gonadotrophin-releasing hormone is well recognized. When it is associated with anosmia due to agenesis of the olfactory bulbs, it is called Kallmann's syndrome. Pituitary deficiency of luteinizing hormone has been described in males. There is little or no testosterone production by the testes, and the seminiferous tubules are immature. However, full maturation of both secondary sexual characteristics and of the seminiferous tubules can be achieved by means of human chorionic gonadotrophin injections, hence accounting for the alternative title of the fertile eunuch syndrome.

In primary gonadal disorders leading to hypogonadism, the characteristic biochemical feature is high serum gonadotrophins. Anorchia is a rare congenital cause in males. More usual causes are trauma, tuberculous orchitis or oophoritis and radiation damage to the gonads. Rarely, there may be autoimmune ovarian failure; this is usually associated with other autoimmune disease such as Addison's disease. Boys with Klinefelter's syndrome (XXY) are usually tall and have abnormally long limbs before puberty but, if testosterone levels are low in the middle to late teens, an element of hypogonadism is added (see below).

Either adenoma or hyperplasia of the pituitary acidophil cells (somatotrophs) can lead to the excessive secretion of GH and the rare condition of ‘gigantism’ analogous to acromegaly in the adult. Sometimes, the adenoma may be of chromophobe cells or of mixed cell type. GH-secreting tumours also occur in multiple endocrine neoplasia, neurofibromatosis and tuberous sclerosis. If the condition arises before puberty, the main feature is excessive growth of the long bones, resulting in a wide span and an increased lower segment to upper segment ratio. Heights of over 2.4 m have been recorded. If hypogonadism does not supervene, the epiphyses may fuse, and acromegalic features may begin to develop. Acromegaly means enlargement
of the extremities; the bones of the hands and feet become broadened and the soft tissues become thicker, so that it is difficult for the patient to get shoes and gloves to fit. The facial appearance gradually changes, the skin and subcutaneous tissues becoming thicker, so that the skin folds are more prominent and the nose enlarged, and the general impression is one of coarsened features. The frontal air sinuses enlarge, so the supraorbital ridges are more prominent. The lower jaw elongates and causes prognathism; dental malocclusion may become a problem. Radiographs may show enlargement of the pituitary fossa, but by no means always. Characteristic tufting of the terminal phalanges and widening of the joint spaces may be seen on hand radiographs. Increased heel pad thickness is usually demonstrated on radiographs. If it is above 23 mm, acromegaly is possibly present; if above 27 mm, acromegaly is certain. Enlargement of the tongue, lips and ears occurs. Thickening of the vocal cords leads to deepening of the voice. Overgrowth of soft tissues at the wrist causes carpal tunnel syndrome in 35 per cent of cases. Evidence of a proximal myopathy is found in one-third of patients. Persistent sweating is a common problem, and girls may be troubled by hirsutism. Persistent headaches caused by stretching of the diaphragma sellae by the enlarging adenoma are a feature in one-third of the patients. Other local pressure effects include visual field loss due to the tumour encroaching upon the optic chiasm. A very early feature may be loss of central vision for red objects; subsequently, the classic bitemporal hemianopia develops, and this may lead on to optic atrophy and total blindness. All organs enlarge, particularly the heart and liver. A cardiomyopathy may be present, and hypertension is a common feature. Frank diabetes mellitus is present in 10 per cent of cases, and a further 33 per cent have impaired tolerance to oral glucose. Goitre may be present, and thyrotoxicosis occurs in 3 per cent of patients. Galactorrhoea due to excessive prolactin secretion may be a feature. In long-standing cases, hypopituitarism develops in about 50 per cent of patients. A resting GH in the fasting state of more than 10 mU/l (5 ng/ml) that does not suppress to less than 2 mU/l (1 ng/ml) after 75 g of oral glucose is diagnostic of acromegaly or ‘gigantism’. Most patients have much higher fasting levels than this, and they may even show a rise in GH levels following glucose. Serum IGF-I levels are also elevated.

**CHROMOSOMAL ABNORMALITIES**

The most commonly recognized chromosomal anomaly leading to tall stature is **Klinefelter’s syndrome**. This syndrome, occurring once in every 500 live male births, is caused by an extra X chromosome, giving a karyotype of 47,XXY. The patient is tall and has eunuchoid proportions (Fig. S.59). The tallness may not be particularly apparent in childhood, but it becomes noticeable during the teens because the hypogonadism delays epiphyseal fusion. The testes are small and firm, and histology shows tubular sclerosis and hyalinization. Variable numbers of Leydig cells may be present, and this accounts for the fact that the serum testosterone may be frankly low or be in the low normal range, follicle-stimulating hormone (FSH) is elevated and luteinizing hormone (LH) usually so, but not if the testosterone is normal. There is a female escutcheon of pubic hair, and facial and body hair is usually diminished. Gynaecomastia is a usual finding. Infertility is the rule, but occasional exceptions have been described. Another syndrome – the supermale, with a karyotype of 47,XYy – also occurs, with a frequency of 1 per 1000 live male births. These individuals are usually very tall. The testes may be normal or small; in the latter case, the testosterone level is low, and FSH and LH are increased. **Figure S.59** Klinefelter’s syndrome. The arms and legs are long in proportion to the trunk, and there is gynaecomastia.
LH are elevated. In some cases, mental disability and a tendency to aggressive behaviour are noted.

The superfemale (47, XXX) tends to have long legs and long fingers, but with a relatively normal span. The face is long and narrow, there is sometimes amenorrhoea, and the secondary sexual characteristics may be poorly developed. Mild mental subnormality is a feature.

MISCELLANEOUS CAUSES

There is a group of miscellaneous causes of tall stature in children of which Marfan’s syndrome is the most commonly recognized (Fig. S.60). This is an autosomal dominant disorder of collagen metabolism. The child is usually within the normal height range for his or her age group but may be inappropriately tall for the family. The extremities are long and thin, and the fingers and toes are spider-like (arachnodactyly). The joints are hyperextensible. There may be dilatation of the aorta, aortic regurgitation, dislocation of the lenses of the eye, a high arched palate and long patellar ligaments. Although the skeletal proportions appear ‘eunuchoidal’, bone maturation, in fact, proceeds at a normal rate. From a radiograph of the hands, the metacarpal index can be calculated by dividing the mean of the lengths by the mean of the widths. An index of more than 8.4 indicates arachnodactyly.

Patients with homocystinuria are rather similar in appearance to those with Marfan’s syndrome. The condition is an autosomal recessive disease. Additional features are mental retardation, a tendency to spontaneous thromboses in arteries and veins, and the presence of homocysteine in the urine. Cerebral gigantism (Sotos syndrome) is characterized by accelerated growth in the first few years of life, although puberty is usually early and causes premature closure of the epiphyses such that a normal adult stature is achieved. The hands and feet are large, and the appearance is somewhat acromegalic, although GH and IGF levels are normal. Mental subnormality is common. No cause has been identified.

Some children who may have suffered brain damage and mental retardation from birth trauma become excessively tall. The cause is unknown.

Lipodystrophy is a rare condition with absent fatty tissue, hyperlipidaemia and diabetes; there may be increased GH levels, and tall stature is a feature. Beckwith–Wiedemann syndrome is rare. It consists of accelerated fetal growth, enlarged tongue, omphalocele, renal medullary hyperplasia and neonatal hypoglycaemia secondary to islet cell hyperplasia and raised insulin levels. Early epiphyseal fusion occurs, and therefore there is no increase in adult height. This syndrome may be associated with disordered regulation of the IGF-II gene transcription.

Failure to enter puberty and complete sexual maturation may result in sustained growth during adult life. The result is a tall stature with eunuchoid appearances. This has been described in conditions where mutation of the oestrogen receptor has taken place, or in cases of aromatase deficiency (enzymatic conversion of androstenedione to oestrogen in extraglandular sites). These abnormalities emphasize the role of oestrogen in epiphyseal closure and termination of skeletal growth.

STOMACH, DILATATION OF

Harold Ellis

(See also ABDOMINAL SWELLINGS, p. 11.)

Dilatation of the stomach presents itself clinically under two totally different aspects: (i) acute; and (ii) chronic.

ACUTE DILATATION OF THE STOMACH

This is generally a serious, but fortunately rare, complication or even a fatal catastrophe arising in the course of some other condition, especially after operations (notably laparotomy), or after abdominal injury.

The diagnosis is generally easy. The abdomen is distended and tympanic; there is constant effort to bring up wind, sometimes in vain, sometimes

Figure S.60 A young male with Marfan’s syndrome.
with copious and recurrent eructations, often with intractable hiccoughs. Sometimes, immense quantities of blackish-brown or greenish-brown fluid flow effortlessly from the mouth and nostrils. The dilatation itself is of the nature of an acute paralysis of the stomach. Diagnosis is confirmed, and indeed treatment initiated, by the passage of a stomach tube that deflates the gastric dilatation.

**CHRONIC DILATATION OF THE STOMACH**

This is due to conditions that cause stenosis at, or more commonly on either side of, the pylorus.

**Causes of stenosis**

These include the following:

- Peptic ulcer, particularly of the duodenum, although much less commonly a benign gastric ulcer at the pylorus or in the antrum
- Carcinoma of the pylorus or antrum
- Other tumours in this region, including leiomyoma, leiomyosarcoma, infiltration with Hodgkin’s disease or lymphosarcoma, or invasion from an adjacent carcinoma of the pancreas or gallbladder
- Congenital pyloric septum
- Adult hypertrophy of the pylorus
- Heterotopic pancreatic tissue
- Adhesions of the duodenum to the liver bed following cholecystectomy

The history in an established case of pyloric stenosis may be absolutely typical. In the case of a peptic ulcer, there may be a long preceding story of ulcer pain. Vomiting is an important symptom, and it occurs in at least nine out of every ten patients. Typically, copious amounts of vomitus are produced in a projectile manner, and the patient will recall (but often only on direct questioning) that he or she has noticed fragments of food, particularly vegetable or fruit debris, that had been ingested one day and vomited up the next or even 2–3 days later. There is really no condition other than obstruction to the gastric outlet in which this state of affairs obtains. Obstruction due to carcinoma, in contrast, often has a shorter history, perhaps of only a few months, and pain is completely absent in about one-third of patients.

**Examination of the patient** often reveals features of importance. There may be evidence of dehydration and loss of weight; indeed, the classic ‘ulcer facies’ applies only rarely to uncomplicated examples of peptic ulcer but is perfectly mirrored in the usual appearance of the victim of long-standing stenosis. A gastric splash that is present 3–4 hours after a meal or drink is elicited in two-thirds of patients with benign stenosis. Often, the patient – when asked directly – will agree that they have noticed a splashing sound when walking or moving about. Visible gastric peristalsis, passing from left to right, is present much less frequently, and still less often the loaded and hypertrophied stomach may actually be palpable as well as audible and visible (see PERISTALSIS, VISIBLE, p. 512). About half of the patients with malignant obstruction will reveal a palpable tumour at the pylorus. Such a mass may, it is true, be felt rarely in the benign case, when a large inflammatory mass is present around the first part of the duodenum. Because of the more rapid progression in the malignant case, gross dilatation of the stomach is much less often seen than in benign obstruction, so that gastric splash and visible peristalsis may not be elicited.

**Radiological investigation** in these cases is mandatory. The findings can be divided into two groups: the first group confirms the presence of an obstruction at the gastric outlet; and the second indicates its pathology. A plain radiograph of the abdomen may itself be at least suggestive of pyloric stenosis by demonstrating a large gastric gas bubble with considerable quantities of retained food particles, as indicated by patchy, translucent areas. A sign of obstruction at the gastric outlet on the barium meal is the large residue of food within the stomach shown after taking a few mouthfuls of barium. Instead of the normal appearance of the barium running down the lesser curvature, the particles of barium can be seen to sink through a layer of fluid and then to rest at the bottom of the greater curve, like a saucer. In the erect position, three layers can be seen: the air bubble above; then the layer of gastric juice; and finally the lowermost layer of barium (Fig. S.61). In the early phase of pyloric stenosis, giant peristaltic waves may be seen passing along the gastric wall; in late decompensated obstruction, the stomach is a large atonic bag. Obstruction of the gastric outlet is confirmed by taking further films at 4–6 hours, when it will be seen that a large residue of barium remains in the stomach (Fig. S.62). In normal circumstances, the stomach is all but empty at the end of 2 hours. It is not always easy to tell the exact cause of the pyloric obstruction. Radiological evidence that a duodenal ulcer is responsible is given by the presence of an active ulcer crater or severe scarring in the duodenal cap. If the obstruction is situated in the antrum of the stomach, it is most probable that the diagnosis is cancer (Fig. S.63), but occasionally a similar appearance is given by a penetrating benign gastric ulcer. A further sign of duodenal bulb obstruction that has been found useful is abnormal dilatability of...
the pyloric canal, which may be seen on screening to dilate up to 2.5 cm or more in width, and then to contract down again to its usual size proximal to the point of stenosis.

Gastroscopy by means of a fiberoptic endoscope may visualize the obstructing ulcer and allow a biopsy to be performed, but adherent gastric contents often obscure the view. This can be obviated by careful pre-examination gastric lavage.

The less common causes of pyloric obstruction mentioned above that may be associated with chronic dilatation of the stomach are rarely diagnosed before laparotomy. However, since obstruction of the gastric outlet almost invariably requires surgical intervention, elucidation of the exact cause preceding operation is a luxury rather than a necessity for the experienced surgeon.

**STOOLS, MUCUS IN**

Harold Ellis

Mucus in the stools is not pathognomonic. It occurs in malignant disease of the colon as a clear glairy substance, often blood-stained, and it has the same character in intususception. The obstruction in both of these conditions accounts for the absence of faecal colouring. Large amounts of mucus may be secreted by extensive benign papillomatous tumours of the colon and rectum. Since this material is rich in
potassium, profound potassium depletion may occur in this condition, leading to weakness, paraesthesiae and even paralysis and vascular collapse. The volume of fluid passed may amount to 2–3 litres daily.

Mucus is often seen with constipated motions, the hard faeces having led to irritation of the large bowel, with consequent increased secretion of mucus as a defensive mechanism against misguided therapy, especially colonic lavage. In severe cases, a motion may consist almost entirely of coagulated shreds, with little faecal matter. In other cases, complete casts of the bowel formed from coagulated mucus are passed; these may be 30 cm or more in length. They may have become broken into fragments, which the patient describes as ‘skins’, which look not unlike segments of tape worm, for which indeed they are, on inadequate examination, easily mistaken.

Patients passing this variety of mucus are said to have membranous or spastic colitis; this is an incorrect term, as no inflammatory process occurs. The term ‘irritable bowel syndrome’ (IBS) is also used for this disorder, which is characterized by colonic abdominal pain, abnormal stools and an alteration in bowel habits. It is more common in females aged 15 to 45 years, but it may occur in either sex under conditions of emotional tension. On examination, the patients often appear anxious and tense and perspire excessively. Curiously enough, this hypersecretion of mucus has almost disappeared during the past 40 years, although the general symptomatology is still recognized. It may be added that the popular treatment of lavage to remove the mucus will be responsible for its continued secretion as a protest against irritation of the mucosa. In the more acute varieties of inflammation of the bowel, the mucus passed is jelly-like and semi-liquid, of varying colour according to the amount of faecal staining. In polyposis coli and severe cases of ulcerative colitis, Crohn’s colitis, enteritis and dysentery, the motions consist of nothing but mucus and blood. Differentiation between the numerous varieties of enteritis and colitis cannot be made upon the basis of the mucus in the stools alone.

**STOOLS, PUS IN**

Harold Ellis

Pus in the stools in sufficient amounts to be recognizable by the naked eye indicates the rupture of an abscess into the intestinal tract. Such recognition is, however, unusual, as even when a large appendicular abscess perforates into the caecum, the pus becomes indistinguishable either from admixture with the faeces (the patient believing they simply have diarrhoea), or on account of its digestion and decomposition. The less the pus is mixed with other intestinal contents, the nearer to the anus must the site of rupture have been. However, the diagnosis of the source of the abscess needs to be determined on other grounds, particularly the history and the results of examination, including that of the rectum and vagina. Abscesses most apt to cause a discharge of pus with the stools are of the appendicular, pericolic, pelvic or other local peritoneal types; of prostatic or perirectal origin; or a pyosalpinx.

Microscopical quantities of pus in the stools may be due to any of the causes already mentioned and, in addition, to affections of the mucous membrane itself. These comprise acute or chronic ulcerative colitis, Crohn’s colitis, dysentery, cholera, dengue, malignant, tuberculous, typhoidal, carcinomatous or venereal ulceration of the bowel. The pus cells may be recognizable as such under the microscope. Examination with the sigmoidoscope, followed by a barium enema X-ray and/or colonoscopy, is invaluable in deciding the diagnosis.

**STRANGURY**

**Ben Challacombe**

Strangury is a collection of symptoms consisting of pain on micturition, pain prior to and after voiding, and severe frequency and urgency. Voiding provides little relief to sufferers, and small volumes, often only drops, are voided on each occasion. The pain is felt in the suprapubic area, and radiates to the perineum and the tip of the penis in males. The most common cause worldwide is a bladder stone repeatedly coming into contact with the bladder neck.

Causes can be divided into the following groups, with the most common pathologies highlighted in italic:

- Psychological: acute anxiety
- Obstruction to urinary outflow
  - Urethral stricture
  - Benign prostatic hyperplasia
  - Prostate carcinoma
  - Gynaecological: gravid uterus, uterine fibroid, ovarian cyst, ovarian carcinoma, uterine prolapse (severe)
  - Impacted urethral calculus/bladder calculus
  - Sexually transmitted infections: gonorrhoea, non-specific urethritis, urethritis
  - Periprostatic or ischiorectal abscess
- Pathology of the bladder wall
  - Lower ureteric calculus
  - Trauma
• Cystitis: acute, recurrent, tuberculous, interstitial cystitis
• Bladder tumour
• Schistosomiasis
• Infiltration by carcinoma of the uterus/rectum

- Pharmacological: medicinal or recreational (ketamine)
- Reflex from local stimulants: painful haemorrhoids

The pain is attributed to irritation of the urothelium (the epithelium lining the urinary tract) and subsequent spasm of the muscles. Many of these conditions cause frequency of micturition (see p. 415) in their own right and are covered in more detail in that section.

A new cause of strangury is ketamine hydrochloride, an anaesthetic agent used in human and veterinary procedures. It has increasingly been used as a recreational drug in either powder or liquid form, which can be sniffed or injected, causing powerful hallucinations. It is a new cause of symptoms of strangury consisting of severe frequency, urgency and dysuria, leading to a small-capacity bladder with severe ulceration and bladder wall thickening.

**STRIAE ATROPHICAe**

Barry Monk

Striae or ‘stretch marks’ are unsightly linear marks due to disruption of the dermal support tissue (e.g. collagen and elastic fibres). Although initially reddish-purple, they later fade to an opalescent whitish colour. They commonly occur following rapid distension, for example during the growth phase of adolescence (lumbosacral region in boys (Fig. S.64), and thighs, buttoks and breasts in girls) and during pregnancy (breasts and abdomen). They are also caused by corticosteroids, which reduce the bulk of dermal support tissue (e.g. Cushing’s syndrome; seen in the flexures), and after steroid therapy, both systemic and topical.

Striae are usually only a cosmetic problem but can, if extensive, ulcerate, particularly following trauma. The sudden development of striae in muscular male subjects should raise the suspicion of the use of anabolic steroids as an aid to body-building, a suggestion that may be vigorously denied by the patient. The co-existence of nodulocystic acne should increase one’s level of suspicion. Caution must be applied in confronting patients with the diagnosis as anabolic steroids may also be associated with problems of aggressive behaviour!

**STRIDOR**

Alex West

(See also WHEEZE, p. 761.)

Stridor is a harsh noise produced during breathing, typically heard as a high-pitched inspiratory wheeze audible at a distance. It is important to note that this sign may indicate an impending life threatening condition so should be investigated urgently, for example in anaphylaxis causing angioedema. It is commonly caused by obstruction affecting the larynx or extrathoracic trachea. The intensity of sound is accentuated by inspiratory negative pressure tending to collapse the extrathoracic airways. Stridor may not be evident on quiet breathing, and is best heard after exercise or during hyperventilation through the open mouth. On auscultation, a fixed inspiratory wheeze is heard over the trachea, becoming fainter over the lungs. When narrowing affects the portion of the trachea within the thorax, the carina or main bronchi, stridor tends to be louder on expiration because the intrathoracic airways partially collapse due to the expiratory rise in intrathoracic pressure. In this situation, it may be difficult to differentiate stridor from the abnormal noisy breath sounds commonly present on quiet breathing in widespread airflow obstruction. These differences, and also the distinction between main airway narrowing and diffuse distal airway narrowing, as in asthma or chronic obstructive pulmonary disease, may be demonstrated by flow–volume loops (Fig. S.65).

Stridor is generated by vibration of the walls of critically narrowed airways, and the pitch is dependent upon the speed of airflow and the density of the inspired air. A low-density gas mixture (helium and oxygen) produces much less turbulence and stridor.
than conventional nitrogen and oxygen mixtures, and therefore may provide useful temporary relief of symptoms in critically breathless patients. With extreme degrees of tracheal narrowing, violent ineffectual respiratory effort, cyanosis and sudden death may result when a relatively minor additional insult, such as a mucus plug, occludes the narrowed trachea.

In children, the onset of stridor is usually alarmingly sudden and due to acute infective conditions (Box S.14); in adults, the onset tends to be insidious and may be confused with late-onset bronchial asthma, chronic bronchitis or other chronic pulmonary disorders (Box S.15).

In many patients, the cause of stridor is obvious, for example when it is caused by secretions lodging in the larynx or trachea in seriously ill patients, or in patients in coma from any cause. In others, local examination of the upper respiratory tract and larynx will provide the diagnosis without much difficulty.

Fibreoptic laryngoscopy and bronchoscopy will often be diagnostic. The greatest difficulty is likely to be encountered in children, in whom the possibility of stridor of acute onset being due to diphtheria should never be forgotten and, in less acute cases, retropharyngeal abscess must be borne in mind. Foreign bodies in the larynx or the main air passages are also a possible source of difficulty in children who are often too young to provide a clear history.

Inspiratory stridor is common in early childhood. The most frequent cause is acute viral laryngotracheobronchitis (croup), but it is essential to consider the full differential diagnosis if tragic and preventable death from acute airway obstruction is to be avoided (Box S.14).

**Box S.14 Causes of acute inspiratory stridor in infants and young children**

<table>
<thead>
<tr>
<th>Congenital</th>
<th>Inhaled foreign body</th>
<th>Croup syndrome</th>
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<tr>
<td>Choanal atresia in newborn</td>
<td>Small plastic toys in crawling children</td>
<td>Diphtheria – any age</td>
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<tr>
<td>Congenital laryngeal paralysis in newborn</td>
<td>Peanuts in older children</td>
<td>Acute laryngotracheobronchitis – usually in 2-year-olds</td>
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<td></td>
<td></td>
<td>Acute epiglottitis – usually in 2- to 7-year-olds</td>
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<td></td>
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<td>Acute pseudo-membranous croup</td>
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<td>Upper airway burns</td>
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<td>Angioneurotic oedema</td>
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<td></td>
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<td>Infectious mononucleosis</td>
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</table>

**Figure S.65** Maximum flow–volume curves represented diagrammatically for: (a) a healthy subject; (b) patients with intrapulmonary airflow obstruction (chronic obstructive pulmonary disease), or (c) patients with obstruction of the upper (extrathoracic) trachea (fixed upper airway). In extrathoracic tracheal obstruction, the reduction in inspiratory flow is equal to or greater than the reduction in expiratory flow, whereas in intrapulmonary airways obstruction, maximum expiratory flow is reduced more than maximum inspiratory flow. TLC, total lung capacity; RV, residual lung volume; PEF, peak expiratory flow.
Acute laryngotracheobronchitis is the common cause of croup in the second year of life. Males are affected most frequently. The main pathogens are parainfluenza virus (50 per cent), respiratory syncytial virus, influenza A and B, rhinovirus, adenovirus and measles. It is usually preceded for 1–2 days by an upper respiratory tract infection before developing a harsh, barking cough and hoarse voice due to inflammatory swelling of the subglottic region. With more severe airway obstruction, chest wall recession occurs, and stridor may be inspiratory and expiratory. Fever, restlessness, hypoxia, tachypnoea, tachycardia and eventually cyanosis herald the risk of sudden collapse and death.

*Acute epiglottitis* is often mistaken for acute laryngotracheobronchitis, but its pathology and clinical course are quite different. The onset is usually rapid and the patient pyrexial and lethargic, complaining of a sore throat. Within a short time, upper airway obstruction becomes obvious, with a soft inspiratory stridor and expiratory sound resembling a snore. The child prefers to sit upright and mouth-breathe, and often drools saliva. Acute epiglottitis is an acute life-threatening form of laryngeal obstruction due to oedema and hyperaemia of the epiglottis, aryepiglottic folds and hypopharynx that results in considerable oedema around the laryngeal inlet, obstructing respiration and swallowing. The changes do not extend below the vocal cords. Septicaemia is invariably present, the usual organism being *Haemophilus influenzae* type B. The clinical features are similar to those of acute obstructive laryngotracheobronchitis, except that there is rapid onset of dysphagia and drooling saliva as well as respiratory distress with stridor and a thick muffled voice.

The child is toxic, and complete respiratory obstruction may supervene very quickly. No attempt should be made to examine the pharynx with a tongue depressor as this may precipitate complete obstruction. Because of the risk of laryngeal obstruction, immediate transfer of the child to a hospital with appropriate facilities is mandatory.
should be arranged for full assessment with a view to nasal or oral intubation and appropriate antibiotic treatment.

**SUCCUSSION SOUNDS**

**Simon Anderson**

Succussion sounds may be detected when a viscus or cavity that contains both liquid and air or gas is shaken. The sounds are detectable with a stethoscope but can be loud enough to be audible at a distance from the patient. Gastric succussion sounds are not necessarily evidence of abnormality; they merely indicate that the viscus contains liquid and gas (e.g. after swallowing a quantity of liquid). Succussion sounds may be heard in the chest in cases of hydropneumothorax or, occasionally, a pyopneumothorax or a haemopneumothorax.

A list of all possible causes of succussion sounds is provided in Box S.16.

**SUCCUSSION SOUNDS IN THE CHEST**

It is almost unknown for a tuberculous cavity to give succussion sounds. Should it do so, the situation would be subapical rather than basal, and thus distinguishable from most cases of hydro- or pyopneumothorax. Hydro- and pyopneumopericardium are also very rare: they are identified by the churning sounds made by the heart beating within the mixture of air and liquid. The cause is generally a tumour of the oesophagus or bronchus extending into the pericardium from behind, a swallowed foreign body such as a fish bone penetrating through from the oesophagus, a subdiaphragmatic abscess penetrating through the diaphragm into the pericardium, or infection of the pericardial sac by a gas-producing microorganism.

A **subdiaphragmatic abscess** containing air due to communication with a hole in a gastric or duodenal ulcer may elevate the diaphragm so high that the condition may be mistaken for hydro- or pyopneumothorax. A decision may be impossible until the position of the diaphragm is ascertained by X-rays and ultrasonography. When the pathology is subdiaphragmatic, the tendency is to displace the heart upwards rather than towards the opposite side of the chest; the contrary is usual in the case of pneumothorax.

**Diaphragmatic hernias**, if large and if the stomach is herniated into the thorax, will show the effect of eating and drinking upon the physical signs, and may point to the diagnosis. X-rays will demonstrate the condition on barium meal.

Most cases of hydropneumothorax present little difficulty in diagnosis, although it may not be easy to ascertain the cause of the condition. If the onset has been sudden, with acute pain in the affected side of the chest, cyanosis and dyspnoea, the most likely cause is tuberculosis. In some instances, an injury or a ruptured emphysematous bulla may have been responsible, but injury seldom produces hydropneumothorax unless a tuberculous or other lesion in the lung was present at the time of the accident. Hydropneumothorax may result from paracentesis thoracis; if bleeding occurs during the puncture, haemopneumothorax will be produced. This is also common after bullet wounds of the chest. Either a hydro- or a haemopneumothorax may become infected with pyogenic organisms and converted into a pyopneumothorax. This may develop from direct extension from an infection in the bronchi (obstruction of a bronchus by foreign body or tumour, bronchopneumonia), or externally (gunshot or other wounds of the chest). Fluid often collects in the pleural cavity when an artificial (therapeutic) pneumothorax has been induced, giving succussion sounds.

**SUCCUSSION SOUNDS IN THE ABDOMEN**

Abdominal succussion sounds are generally produced from slow-growing or chronic conditions.

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**Box S.16  Causes of succussion sounds**

**In the thorax**
- Hydropneumothorax
- Pyopneumothorax
- Haemopneumothorax
- Diaphragmatic hernia
- Subdiaphragmatic abscess communicating with the stomach or duodenum, or infected with Escherichia coli; in either case, gas and pus are present
- Hydropericardium
- Pyopneumopericardium

**In the abdomen**
- The normal stomach
- Gastric outlet obstruction e.g. pyloric stenosis, stomach cancer
- Gross dilatation of the caecum
- Pneumoperitoneum due to:
  - Perforated gastric ulcer
  - Perforated duodenal ulcer
  - Perforated typhoid ulcer of the intestine
  - Perforated carcinoma of the colon
  - Production of gas by E. coli, either in a local abscess (e.g. appendicular or subdiaphragmatic) or in the general peritoneum
- Subdiaphragmatic abscess communicating with the interior of the stomach
- Air and urine in the bladder (see PNEUMATURIA, p. 519)
- Gas production by E. coli in a large pyonephrosis, infection by a gas-producing microorganism of an ovarian cyst or other collection of fluid

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Hydro- and pyopneumopericardium are also very rare: they are identified by the churning sounds made by the heart beating within the mixture of air and liquid. The cause is generally a tumour of the oesophagus or bronchus extending into the pericardium from behind, a swallowed foreign body such as a fish bone penetrating through from the oesophagus, a subdiaphragmatic abscess penetrating through the diaphragm into the pericardium, or infection of the pericardial sac by a gas-producing microorganism.

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SWEATING, ABNORMALITIES OF

Sweating is the normal response to exercise or excessive heat, although there is marked physiological variation between individuals. Fever and sweating accompany infections, and drenching sweats usually occur, with a fall of temperature. Sweat regulation can be unbalanced by pyrogens, and sweating may occur out of phase with the fever, for example in some cases of tuberculosis, brucellosis, HIV and lymphoma. These conditions are often associated with night sweats. Generalized hyperhidrosis may also be produced by drugs (e.g. alcohol, pilocarpine and tricyclic antidepressants). Hyperhidrosis may cause maceration of the skin, increasing the risk of fungal or pyogenic skin infection.

A ‘cold and clammy skin’, where sweating is associated with cutaneous vasoconstriction, occurs in hypoglycaemia, the dumping syndrome, alcohol and drug withdrawal, shock and syncopal states, and also in intense pain. Increased sweating is seen in many endocrinological disturbances – hyperthyroidism, hyperpituitarism and acromegaly, for example. It has also been described with phaeochromocytoma, carcinoid syndrome and gout. Sweating is a feature of rickets, pink disease and infantile scurvy (Barlow’s disease).

LOCAL HYPERHIDROSIS

Local increase in sweat production is seen with organic neurological lesions (e.g. brain tumours and spinal cord injuries), and can help to localize the neurological defect. Local hyperhidrosis of the palms, soles and/or axillae is not uncommon, and this occurs in patients who are otherwise perfectly normal; the sweating increases with embarrassment, anxiety and during the summer months. Sweat can literally drip from the hands, making paperwork extremely difficult. The keratin of the soles becomes macerated, and secondary infection causes an offensive odour (osmidrosis). In pachydermoperiostosis, local hyperhidrosis occurs over the skin folds of forehead and extremities.

Granulosis rubra nasi is a rare, genetically determined disease in which profuse sweating of the tip of the nose is associated with a diffuse erythema and the formation of minute, dark red papules. The disorder usually begins in early childhood and subsides

**Box S.17 Hyperhidrosis**

<table>
<thead>
<tr>
<th>Generalized</th>
<th>Local</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Exercise</td>
<td>• Infantile scurvy</td>
</tr>
<tr>
<td>• Raised ambient temperature</td>
<td>• Pink disease</td>
</tr>
<tr>
<td>• Anxiety</td>
<td>• Hyperthyroidism</td>
</tr>
<tr>
<td>• Infections/pyrogens</td>
<td>• Hyperpituitarism</td>
</tr>
<tr>
<td>• Drugs: alcohol, pilocarpines, Granulosis rubra nasi tricyclic antidepressants</td>
<td>• Acromegaly</td>
</tr>
<tr>
<td>• Hypoglycaemia</td>
<td>• Phaeochromocytoma</td>
</tr>
<tr>
<td>• Dumping syndrome</td>
<td>• Carcinoid</td>
</tr>
<tr>
<td>• Alcohol/drug withdrawal</td>
<td>• Gout</td>
</tr>
<tr>
<td>• Shock/syncope</td>
<td></td>
</tr>
</tbody>
</table>
at puberty. It must be distinguished from rosacea, lupus erythematosus and lupus vulgaris.

**Miliaria**
Miliaria are lesions caused by blockage and rupture of the sweat ducts; they are most often seen in tropical conditions of heat and high humidity, which are simulated in the UK by polythene occlusion of the skin, or in neonatal nurseries. There are three forms depending on the depth of duct obstruction:

- **Miliaria crystallina** (sudamina): superficial blockage causes ‘crystal’-clear vesicles just below the epidermal surface, with little inflammatory reaction and few symptoms. These are often seen on the trunk during febrile illnesses.
- **Miliaria rubra** (prickly heat): the rupture occurs in the mid-epidermis, resulting in tiny, intensely pricking red papules. These may erupt widely in persons recently arrived in tropical conditions, or they may be confined to friction areas and flexures.
- **Miliaria profunda** (mammillaria): the blockage occurs within the dermis. Lesions are easily overlooked as they are neither red nor uncomfortable but appear as firm papules (1–3 mm) on the body or limbs. They may follow repeated attacks of miliaria rubra. The natural history of miliaria depends on environmental factors; if sweating continues, the lesions continue to erupt but, if sweating is arrested (e.g. by entering an air-conditioned or cool area), healing can commence.

**Anhidrosis**
Anhidrosis is much less common than hyperhidrosis; sweating may be either diminished or totally suppressed, and either the whole skin or only some particular area is affected. A lack of sweat glands may occur with skin dystrophies (e.g. congenital ectodermal dysplasia) or atrophy (e.g. scleroderma). The sweat gland population may also be diminished in ichthyosis and Anderson–Fabry’s disease. In these situations, heat regulation may be impaired. Suppression of sweating is characteristic of heatstroke, although the mechanism of heat stress and acclimatization is still poorly understood. Organic brain damage at any level, especially of the hypothalamus, can result in complete anhidrosis. Localized anhidrosis can follow spinal cord lesions, for example syringomyelia or the neuropathy of diabetes or leprosy. It can thus help in localizing neurological lesions. Sympathectomy abolishes sweating. Generalized decreased sweating and dry skin is a well-known sign in myxoedema. Blockage of the sweat ducts can occur in atopic eczema, psoriasis and lichen planus, and it can be associated with crises of itching, as in miliaria rubra.

**Osmidrosis**
Osmidrosis is foul-smelling sweat. Sweat is usually odourless, but various substances may be excreted in sweat, including garlic, drugs (e.g. dimethyl sulphoxide) and arsenic. In the past, physicians have remarked on the particular odour of sweat: in diabetes, gout, scurvy, typhoid, uraemia (see also SMELL, ABNORMALITIES OF, p. 623). Hyperhidrosis, especially of the soles, is commonly accompanied by a foul odour due to bacterial overgrowth. Imaginary osmidrosis is a well-recognized paranoid delusional symptom. Personal body odour is largely determined by apocrine gland secretion.

**Chromhidrosis**
Chromhidrosis is usually due to coloured sweat. Perhaps 10 per cent of normal people have coloured apocrine sweat (blue, yellow or green), and rarely areas of ectopic apocrine glands can give rise to areas of discoloured skin, where it is possible to see tiny beads of coloured sweat. The pigments are lipofuscins. In pseudochromhidrosis, colourless sweat becomes coloured on the surface of the skin, due usually to chromogenic bacteria. Occasionally, workers in the chemical industry can develop coloured eccrine sweat that can be shown to contain dyes. Certain drugs (e.g. rifampicin) may give rise to coloured sweat.

**Urhidrosis**
The urea content of sweat increases with serum urea concentration. Uraemic sweat has a particular odour and, after evaporation, leaves a visible deposit of urea crystals on the skin. This is known as ‘uraemic frost’.
TACHYPNOEA
Alex West

Tachypnoea signifies an increase in the rate of breathing, as may occur normally during exercise, or abnormally with a number of clinical disturbances, especially those associated with hypoxia. In most clinical situations, tachypnoea is associated with symptoms of breathlessness (see DYSPNOEA, p. 144).

In health, both the rate and depth of respiration are controlled by brainstem respiratory centres to maintain a constant arterial blood oxygen level. The relationship between ventilatory rate and depth, demands for oxygen uptake and requirements for carbon dioxide elimination is controlled by complex interaction between: (i) mechanoreceptors in the airways and lung parenchyma; (ii) peripheral chemoreceptors in the carotid and aortic bodies; and (iii) central chemoreceptors mostly located on or beneath the ventral surface of the medulla. The automatic control system situated in the brainstem is concerned primarily with oxygen, carbon dioxide and acid–base homeostasis. When metabolic requirements are modest, this automatic system can be overridden by the voluntary control system that arises in the cerebral cortex to allow other activities such as talking, coughing and singing to occur. A low arterial partial pressure of oxygen, high arterial partial pressure of carbon dioxide and an acid pH can all stimulate ventilation.

There are many causes of tachypnoea. The increased metabolic demands of fever or circulatory shock from myocardial infarction, haemorrhage, trauma or pulmonary embolism often manifest clinically with rapid breathing. Stimulation of the central chemoreceptors by metabolic acidosis in uncontrolled diabetes mellitus, renal or hepatic failure, or acute circulatory collapse as well as salicylate overdose may all cause tachypnoea. Stimulation of lung stretch receptors with a reflex increase in ventilatory drive may follow acute pulmonary emboli, acute pneumonias, asthma, aspiration of the gastrointestinal contents, pulmonary fibrosis or bullous lung disease, and provoke tachypnoea out of proportion to the hypoxia so commonly associated with these conditions. Neurological causes of tachypnoea include cerebral haemorrhage, meningitis, encephalitis, head injury or primary abnormalities in the brainstem.

Breathing is under voluntary control, and deep inspiration in excess of ventilatory requirements can be taken to anticipate exercise or facilitate talking or singing. Inappropriate hyperventilation is common and may present with complaints of breathlessness, associated with light-headedness, dizziness, peripheral paraesthesia, headache or anxiety. Even tetany may occur if a significant respiratory alkalosis results from over breathing.

Clinical features suggesting a diagnosis of hyperventilation syndrome include complaints of breathlessness at rest for no apparent reason, a paradoxical absence of significant breathlessness on exercise, and marked variability of symptoms. Clinical examination with routine respiratory investigations (including a chest radiograph) may reveal no abnormality, although the occasional patient does exhibit a sighing, irregular pattern of breathing. The most common cause of this condition is anxiety.

TASTE, ABNORMAL PERCEPTION OF
David Werring & Mark Kiniron

Taste perception comprises the basic elements of sweet, salty, bitter and sour. Disorders of taste are much less common than those of smell. In fact, the complaint of abnormal taste is usually found to be due to olfactory dysfunction with preserved taste. This is because, in the overall perception of ‘taste’, smell and taste proper are inextricably linked; a patient with anosmia will complain that food is ‘tasteless’ because the normal tongue can only distinguish the basic tastes of sweet, salt, sour and bitter. Patients with altered or lost taste should therefore be asked about symptoms of head injury or recent upper respiratory illness (which could disturb smell) as well as Bell’s palsy (which may disturb taste). Both taste and smell may be lost in acute upper respiratory infection due to a coated tongue and inflamed nose. Smell and taste must be tested separately in every case. Taste can be tested with solutions of sugar, sodium chloride or quinine applied to each quadrant with a cotton applicator. Cards are used for the patient to point to their impression of taste, and water is used to rinse the mouth between applications.

True loss of taste is found in disease of the tongue itself, the facial nerve (chorda tympani branch), the glossopharyngeal nerve, and the glossopharyngeal nucleus in the medulla. Loss of smell by itself occurs in inflammatory conditions of the nose, in tumours of the anterior fossa (which compress the olfactory nerves), and frequently as a result of fractures of the cribriform
The appreciation of taste is a function of the tongue; fibres from the anterior two-thirds pass via the chorda tympani to the geniculate ganglion, passing from there to thepons by the nervus intermedius. Fibres from the posterior third travel in the glosopharyngeal nerve. In the pons, taste fibres pass into the tractus solitarius, and from the nucleus of this tract a gustatory lemniscus is formed, which, after decussating, passes upwards near the midline to the thalamus, and from there to the gustatory cortex at the bottom of the postcentral gyrus. Disorders of taste can be classified as absence of taste (ageusia), diminished taste (hypogeusia), increased sensitivity to some or all taste qualities (hypergeusia), distortion of taste (parageusia or dysgeusia), or gustatory hallucinations. Persistent foul taste can be considered as a specific form of dysgeusia and termed cacogeusia. Common causes of taste disturbance are listed in Box T.1.

From the diagnostic point of view, impairment of taste sensation is important only when it is persistent or recurrent, and even then it is usually due to a primary condition of the mouth, nose, lung or gastrointestinal tract. In the absence of one of these explanations, diagnosis becomes difficult, and a neurological cause should be considered. The most common neurological cause of loss of taste is Bell's palsy, in which it often occurs as an early and transient feature, the loss being confined to the anterior two-thirds of the tongue on the same side as the facial paralysis. This may be asymptomatic or symptomatic. Recovery of taste within 14 days usually indicates that a full recovery will occur.

A persistent facial palsy and loss of taste may be due to the spread of inflammatory disease from the middle ear. Taste may be impaired due to disease process affecting lower cranial nerves (e.g. carcinomatous meningitis or skull base tumours), but this will usually be accompanied by more obvious symptoms and signs. Organic lesions of the temporal lobe uncus (glioma, demyelination, stroke, arteriovenous malformation or cortical dysplasia) can cause uncinate fits in which there is a hallucination of smell or taste, usually unpleasant, followed by salivation, chewing or sucking movements of the mouth and jaws, and disturbance of consciousness that may be mild or progress to loss of consciousness and a generalized convulsion. The distortions of taste that occur during pregnancy are neither delusional nor hallucinatory, but their origin is not understood. Disturbances of smell or taste are uncommon manifestations of hysteria. Loss of taste and smell occasionally occurs as part of a migraine aura.

### Box T.1 Disorders of taste

<table>
<thead>
<tr>
<th>Ageusia (impairment or loss)</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Local mouth disorders</td>
</tr>
<tr>
<td>- Coated or geographic tongue</td>
</tr>
<tr>
<td>- Epithelioma</td>
</tr>
<tr>
<td>- Glossitis</td>
</tr>
<tr>
<td>- Xerostomia (dry mouth): Sjögren’s syndrome, radiation therapy, ptyalopsia</td>
</tr>
<tr>
<td>• Neurological causes</td>
</tr>
<tr>
<td>- Glossopharyngeal nerve damage (e.g. meningeval infiltration, skull base tumour, carotid dissection)</td>
</tr>
<tr>
<td>- Facial nerve chorda tympani damage (e.g. Bell’s palsy)</td>
</tr>
<tr>
<td>- Thalamic damage (e.g. stroke, tumour)</td>
</tr>
<tr>
<td>- Pontine damage (e.g. stroke, demyelination)</td>
</tr>
<tr>
<td>• Drugs (e.g. penicillin, angiotensin-converting enzyme inhibitors, calcium-channel blockers, nasal sumatriptan)</td>
</tr>
<tr>
<td>• Other causes</td>
</tr>
<tr>
<td>- Diabetes</td>
</tr>
<tr>
<td>- Hypogonadism</td>
</tr>
<tr>
<td>Parageusia/dysgeusia (altered taste)</td>
</tr>
<tr>
<td>• Pregnancy or menstrual cycle changes</td>
</tr>
<tr>
<td>• Glossitis</td>
</tr>
<tr>
<td>• Hysteria</td>
</tr>
<tr>
<td>• Drugs (e.g. metronidazole, corticosteroids, lamotrigine)</td>
</tr>
<tr>
<td>• Cortical damage (e.g. stroke affecting gustatory cortex, rare)</td>
</tr>
<tr>
<td>Gustatory hallucinations</td>
</tr>
<tr>
<td>• Uncinate pathology (e.g. tumour, demyelination)</td>
</tr>
<tr>
<td>• Psychiatric disease</td>
</tr>
<tr>
<td>Cacogeusia (foul taste)</td>
</tr>
<tr>
<td>• Local oral disease: caries, stomatitis, gingivitis, glossitis</td>
</tr>
<tr>
<td>• Gastrointestinal disease: carcinoma, gastritis, pyloric stenosis</td>
</tr>
<tr>
<td>• Lung disease: abscess, tuberculosis, bronchiectasis</td>
</tr>
</tbody>
</table>

### TESTICULAR ATROPHY

Mark Kinirons

Apart from the physiological atrophy of the testes that occurs with advancing age, and may begin as early as the age of 50 years, the causes of testicular atrophy can be classified under the headings hypothalamic/pituitary conditions, testicular conditions, suprarenal disorders, general disease and drugs (Box T.2).

#### HYPOthalamic/PiTUITARY CONDITIONS

The testes contain two main components: the seminiferous tubules, which are responsible for the production of spermatogenesis; and the interstitial tissue, which contains Leydig cells that secrete testosterone. Spermatogenesis is under the control of one gonadotrophin, follicle-stimulating hormone (FSH), and testosterone secretion is stimulated by the other gonadotrophin, luteinizing hormone (LH), both of which are produced by the pituitary gland in response to the
secretion of the hypothalamic hormone gonadotrophin-releasing hormone (GnRH). It follows from this that any hypothalamic or pituitary condition that reduces GnRH or gonadotrophin secretion is likely to lead to testicular atrophy. Probably the most common hypothalamic condition is a congenital lack of GnRH, which, when associated with anosmia, is called Kallmann’s syndrome (see PUBERTY, DELAYED, p. 532). In this condition, there is a midline defect with agenesis of the olfactory bulbs and other midline anomalies such as cleft palate and hare lip. The patients are sexually immature and are tall, with eunuchoidal skeletal proportions, having long arms and legs relative to the trunk. Kallmann’s syndrome is an autosomal dominant condition with variable penetrance, and it occurs in 1 in 10 000 male births and 1 in 50 000 female births.

Isolated LH deficiency is a rare condition causing the ‘fertile eunuch syndrome’. There is little or no testosterone production by the Leydig cells, and the seminiferous tubules are immature because an adequate concentration of intracellular testosterone is necessary for this development and for the production of spermatozoa. The patients are sexually immature and have eunuchoidal skeletal proportions. They can be made sexually mature by treatment with human chorionic gonadotrophin, which has LH-like activity, and this also fully develops the seminiferous tubules.

The Laurence–Moon–Biedl syndrome is an uncommon condition that is a hereditary autosomal recessive disorder with variable penetrance. There are low levels of gonadotrophins, with consequent retarded sexual development, obesity, mental disability, retinitis pigmentosa and either polydactylism or syndactylism.

Raised prolactin levels produced in men by macroprolactinomas of the pituitary or as a consequence of drug therapy (phenothiazines, metoclopramide, haloperidol, pimozide, methyldopa, reserpine or cimetidine) lead to reduced gonadotrophin secretion by the pituitary, and consequently to impotence, infertility and testicular atrophy.

**TESTICULAR CONDITIONS**

Conditions affecting the testes directly are a much more common cause of atrophy than are those secondarily related to hypothalamic or pituitary disorders. Trauma, such as being struck in the testes by a cricket ball or falling astride a fence, may lead to an intratesticular haematoma, which causes pressure necrosis of the seminiferous tubules and eventual atrophy of the testis. Any disturbance of the blood supply to the testes, particularly torsion, can lead to testicular atrophy.

Mumps occurring in males after the age of 13 years is an important cause of testicular atrophy. Permanent atrophy may occur in 36 per cent of cases. A wide range of other infections mentioned in Box T.2 may have the same effect.

Testes that have been situated in the inguinal canal (cryptorchidism) or ectopically in the superficial inguinal pouch, perineum or upper thigh may fail to develop properly and remain small. Cryptorchidism, which is unilateral in 84 per cent of cases, occurs in 3 per 1000 of the adult male population. Cryptorchidism may be associated with Kallmann’s syndrome (see above) but, more importantly, also with Klinefelter’s syndrome (see Fig. S.59). This condition is relatively common, occurring in 1 per 500 male births. It is due to the presence of an extra X chromosome, giving
a karyotype of 47,XXY, although sometimes mosaicism occurs with a karyotype of 46,XY/47,XXY. The patients are tall with eunuchoidal proportions, and they have small, firm testes that, if biopsied, show tubular sclerosis and hyalinization. Variable numbers of Leydig cells are present. There is invariably gynaecomastia, and there is a tendency to low intelligence. The gonadotrophins LH and FSH are increased.

Patients with partial androgen resistance (Reifenstein’s syndrome) may present with small testes, although there is usually also hypospadias and a small penis. The condition is an X-linked recessive with a 46,XY karyotype.

Noonan’s syndrome may present with atrophic testes, although more commonly the testes are undescended.

Dystrophia myotonica (Fig. T.1) is an autosomal dominant condition with a considerable degree of penetrance. Although it commonly appears to present in early middle age, careful studies have shown that evidence of it is usually present in adolescence. Testicular atrophy, cataract and baldness are usual features accompanying the muscular features, which consist of atrophy of the sternomastoids, facial muscles and distal muscles of the limbs, and a characteristic delay in the ability to relax the muscles, exemplified in the prolonged handshake. In addition, the patient usually displays progressive intellectual and psychological impairment.

Haemochromatosis leads to iron deposition in many tissues, including the pituitary and testes. The patient usually has testicular atrophy secondary to low levels of gonadotrophins. The patient has slate-grey skin and may have an enlarged liver and spleen. Feminizing tumours of the ovary or suprarenal gland are extremely rare but may cause atrophy of non-tumourous testicular tissue because of oestrogen secretion. The patient usually also displays gynaecomastia.

SUPRARENAL DISORDERS

Cushing’s syndrome may lead to some degree of testicular atrophy because the high levels of plasma cortisol reduce LH secretion. Similarly, children with congenital suprarenal hyperplasia have small testes, despite development of the penis and secondary sexual hair, because the raised concentrations of androgens suppress the gonadotrophins.

GENERAL DISEASE

Alcohol abuse and cirrhosis of the liver are important causes of testicular atrophy. Alcohol impairs gonadotrophin release from the pituitary, but it also has a direct effect on the testes, lowering testosterone synthesis. Spermatogenesis is also impaired. Testicular atrophy is found in 10–50 per cent of alcoholics without liver disease. Cirrhosis of the liver causes decreased testosterone secretion and an increased conversion of testosterone to oestrogen. This leads to an increase in the carrier protein, sex hormone-binding globulin, which binds more avidly with what testosterone there is, so that the level of free testosterone falls even further. Between 30 and 75 per cent of alcoholics with cirrhosis have testicular atrophy.

Approximately one-third of males with sickle-cell anaemia have testicular atrophy, presumably secondary to impairment of blood supply or infarction of testicular tissue.

Testicular atrophy is frequently found in men with the AIDS. The hormonal changes present in AIDS suggest a non-specific response to systemic illness – low testosterone without an appropriate increase in LH. The disease may involve the testes directly, as virus has been found in lymphocytes in the seminiferous tubules, in the interstitial cells of the testes and within spermatogonia. Opportunistic infections (e.g. cytomegalovirus infection of the testes) have been noted in approximately 30 per cent of cases at post-mortem examination. These patients suffer from hypogonadism.
DRUGS
Drugs used in the treatment of carcinoma of the prostate lead to testicular atrophy. Oestrogen therapy was commonly used in the past, but recently long-acting analogues of GnRH have been used. These act by exhausting the pituitary gonadotroph cells and lowering the levels of gonadotrophins, so that a medical ‘orchidectomy’ occurs.

Chemotherapy, particularly with cyclophosphamide and chlorambucil, leads to testicular damage. Pubertal, but not prepubertal, boys seem most vulnerable. Marijuana has a direct effect on the hypothalamus, pituitary and testis, and spironolactone not only impairs testosterone synthesis but also antagonizes the action of testosterone.

TESTICULAR PAIN
Ben Challacombe
(See also TESTICULAR SWELLING, p 666.)

Pain in the testis of varying degree may be present in many conditions, which may be discussed under separate headings as follows: (i) diseases of the body of the testis or epididymis; (ii) pathology within the coverings of the testicle; (iii) pathology of the spermatic cord; (iv) a retained or misplaced testicle; (v) referred pain from lesions remote from the testis.

DISEASES OF THE BODY OF THE TESTIS OR EPIDIDYMIS
See Box T.3.

Inflammatory lesions
Inflammatory lesions may attack the testis proper or more commonly may begin in the epididymis. The investing tunica vaginalis distends with inflammatory exudate to form a secondary hydrocoele. This tender mass may be mistaken for a swelling of the testis proper, and the condition is frequently labelled an ‘acute epididymo-orchitis’, although more accurately this is usually an ‘acute epididymitis’. An acute epididymitis arises most commonly by retrograde spread of infection from the prostatic urethra via the vas deferens or its accompanying lymphatics. Acute or recurrent epididymitis can sometimes lead to a chronic epididymitis.

Acute epididymitis begins as a painful thickening of the epididymis associated with febrile symptoms. Before any actual pain is noticed in the testis, there is often a sense of discomfort and weight over the external abdominal ring and inguinal canal due to the inflammatory process extending along the vas deferens. The swelling of the epididymis increases, and with it there is a secondary effusion of exudate into the tunica vaginalis (secondary hydrocoele), causing swelling of its body and an increase in pain. The whole organ thus becomes enlarged, and it is often exquisitely tender, the touch of the clothes or the gentlest examination causing pain. The vas deferens and tissues of the spermatic cord are thickened and tender.

Prior to effective antibiotic treatment, by far the most common cause of an acute epididymitis was acute gonorrhoeal urethritis. During the disease, the prostatic portion of the urethra frequently becomes infected, when the orifices of the ejaculatory ducts may share in the infection and inflammation may be conveyed by the vas deferens to the testicle. Infection may also arise following an attack of non-specific urethritis (NSU), acquired as a result of sexual intercourse with a partner suffering from non-specific genital infection.

NSU is today the most common sexually transmitted disease in the Western world, and at least 1 per cent of all male cases develop epididymitis. Chlamydia trachomatis is commonly found and appears to be the cause in around 50 per cent of cases. The gonorrhoeal form of acute epididymitis usually resolves slowly and shows little liability to suppurate, whereas the inflammation resulting from a staphylococcal or streptococcal infection may break down into a testicular abscess.

Acute epididymitis may also arise from septic processes in the urethra following the passage of
a catheter or of instruments, after transurethral prostatic resection, from infection behind a urethral stricture or a calculus in the prostatic urethra, or as a complication of a urinary infection by Escherichia coli or other organisms. It may also follow prostatic massage. The onset of pyrexia with pain and rapid swelling of the testis should lead to suspicion of a urinary tract infection. Bacteriological examination of any urethral discharge and of the urine is essential (see URETHRAL DISCHARGE, p. 722).

In non-specific epididymitis, there may be no evidence of urethral infection, and standard bacteriological studies are entirely negative; the condition sometimes arises after unaccustomed exercise, and it has been attributed to a reflux of urine down the vas. The testicle becomes painful and enlarges rapidly in the same manner as in acute inflammation from urethral infection; under appropriate conservative treatment by means of a scrotal support, this gradually resolves. Less frequently, testicular inflammation may occur after a direct injury to the organ, such as a blow or squeeze.

The pain in an acute inflammation is generally of an aching character at first, felt not only in the testis but also at the external abdominal ring, and often as a heavy dragging pain in the inguinal or iliac areas of the affected side. As the testis enlarges, the local pain becomes more severe, so that the swollen gland is exquisitely tender to pressure or to the touch. After a few days, the pain to a large extent subsides, but it remains as a dull ache until the swelling becomes greatly reduced and usually does not disappear entirely until the organ returns to the normal size. In a few cases, pain may remain and cause some difficulty in the diagnosis from an incipient tuberculous lesion. In other cases, the persistence of the pain and swelling may indicate the formation of an abscess in the testicle, when the swelling increases, the skin covering it becomes reddened and oedematous, and a soft area becomes evident in one aspect of the organ.

Acute orchitis
Acute orchitis may complicate acute specific parotitis (mumps), especially when this occurs in adolescents or adults. The testicular pain and swelling usually occur within a week of the parotid swelling. Both testes may be affected, and the result may be bilateral testicular atrophy, with resultant infertility. Much less often, the testis may be affected in typhoid, scarlet fever or influenza.

Tuberculosis of the testis
Tuberculosis of the testis is still common today in many parts of the world, including India and the Far East, but it is rare in the UK. It usually arises secondary to tuberculous disease of the kidney, bladder, prostate or seminal vesicles. It is most frequently seen in young adults. It begins as a localized deposit in almost all cases, causing a rounded, firm nodule in the epididymis, usually in the lower pole. This nodule may remain unaltered for many months, or it may enlarge, soften, become adherent to the skin and coverings of the testis, or actually ulcerate through them to form a discharging sinus in the scrotum. The small nodule in the epididymis is usually painless at first but later, as it gradually enlarges, it causes an aching pain in the organ. There may be an associated hydrocoele of the tunica vaginalis. Other nodules may be formed in the epididymis, or the body of the testis may become involved, while small shot-like thickenings may commonly be felt in the course of the vas deferens, or a progressively increasing thickening as it is traced down to the epididymis. In the more advanced stages, nodules may be felt upon rectal examination in the seminal vesicles or prostate, or there may be some in the epididymis of the other side.

Tuberculous disease of the testis usually presents some difficulty in its diagnosis from non-specific epididymitis, particularly when it has an acute onset. In an early case, the occurrence of one or more nodules in the epididymis, which are painful on pressure and which have not resulted from a preceding acute epididymitis, should always suggest a tuberculous focus, and a careful search should be made for other tuberculous lesions in the body. The urine is cultured for Mycobacterium tuberculosis, and contrast computed tomography or intravenous pyelography performed. If no evidence of tuberculosis is found, the gradual subsidence of the lesion under careful observation will indicate that it was a non-specific epididymitis. In later stages, the diagnosis is less difficult; the gradual enlargement of the nodules, the infection of the vas or other genitourinary organs with tuberculosis, and the tendency of the focus in the epididymis to soften and become adherent to the scrotal coverings and to produce an indolent sinus are classical features.

Malignant tumours of the testis
Malignant tumours of the testis may give rise to pain in the organ but, as a rule, pain is experienced only in the later stages of the disease. This topic is considered fully in TESTICULAR SWELLING (p. 666).

Torsion of the testis
Torsion of the testis on its vascular pedicle may occur in a testis that has a mesorchium or in one which is ectopic.
It occurs most commonly in infants or in youths soon after puberty; the exciting cause may be some mild exertion or movement such as crossing the legs or turning over in bed. There may be a history of repeated minor attacks before complete torsion takes place, and the other testis may have suffered similar incomplete attacks or be found to be unduly mobile or horizontally placed. At the moment of torsion, there is a severe sickening pain that may at first be felt in the abdomen but is quickly localized to the testis; the boy may even say that his testicle has twisted. There is usually nausea and sometimes vomiting. The testis forms a tense, tender swelling in the upper part of the scrotum or at the external abdominal ring, and the scrotum below is empty. This sign serves to distinguish the condition from a strangulated hernia or an inflamed lymph node.

In acute epididymitis, the testis is in its normal position, and there may be evidence of urethral discharge or of a urinary tract infection. Because of the initial abdominal pain and vomiting, the condition has been mistaken for acute appendicitis, but adherence to the rule of examining the scrotal contents in all abdominal cases should prevent this error. Testicular torsion requires urgent surgical exploration with orchidopexy or orchidectomy to the affected side if it is non-viable.

Cysts of the epididymis
Cysts occur most frequently in connection with the epididymis. These cysts are quite different from hydrocoele of the tunica vaginalis. They cause a swelling of varying degree in the scrotum, and usually an aching in the testicle, groin or lumbar region. These cysts are usually placed above and to the outer side of the testis, occasionally behind it. They move with the organ and can usually be distinguished from the latter by the test of translucency. They may be multiple and are frequently bilateral. Their increase in size is very slow, but they may cause aching pain in the testicle by pressure upon, or stretching of, the tissues of the epididymis. They can be distinguished from hydrocoele of the tunica vaginalis by the position of the swelling relative to the testicle, and by the fact that the fluid contained in them is colourless or slightly opalescent from the contained spermatozoa, in distinction from the straw-coloured clear fluid of a vaginal hydrocoele.

A small cyst on the upper pole of the testis, a few millimetres in diameter (appendix testis) or at the upper end of the epididymis (appendix epididymis) is extremely common. Occasionally, one or other may undergo acute torsion and result in sudden and severe testicular pain that mimics testicular torsion. This may lead to a blue spot sign, where the torted appendage is visible through the scrotal skin as a distinct blue area.

Syphilitic disease of the testis
Syphilitic disease of the testis, once common but now a rarity, causes very little pain in the organ, but there is often a sense of dragging or heaviness. Syphilis may attack the testicle in several different ways, producing diffuse interstitial orchitis or localized gummatous orchitis.

The outstanding feature of syphilitic disease of the testicle is that it affects the body of the testis rather than the epididymis. In the interstitial form, there is thickening of the intertubular connective tissue, forming young connective tissue and yielding fibrous tissue. The subsequent contraction of this fibrous tissue may cause atrophy of the testis. The testis may show small gummas in addition to the diffuse orchitis, or, if the inflammation is more localized, gummas may be the main feature, these varying in size from that of a pea to that of a walnut, or larger. The epididymis is only rarely affected.

In congenital syphilis both the interstitial and gummatous forms exist; they usually occur in childhood or in young adult life, and in many cases the affection is bilateral. Syphilitic inflammation of the testicle may be accompanied in either the acquired or the congenital form by a vaginal hydrocoele. A gummatous testis may ulcerate through the scrotum, usually in front, producing a circular ‘punched-out’ ulcer with a slough in the base.

There is a sense of weight in the scrotum rather than pain, and often an aching or dragging feeling in the inguinal or lumbar region. On palpation, the body of the testis feels enlarged and nodular, but the epididymis can usually be distinguished from the testis and found to be unaffected. Testicular sensation is lost. The tissues of the cord remain unthickened. Tertiary syphilitic lesions of the testis give rise to very little tenderness on palpation.

The diagnosis of syphilitic disease of the testis is usually simple. There may or may not be a history of syphilis, but other signs of the disease should be looked for – thus, in the acquired form, any scar of previous ulceration or periosteal thickening, or, in the congenital variety, signs in the teeth, eyes or ears. Syphilitic disease is distinguished from tuberculous disease of the testis by the fact that the epididymis is usually not involved; the cord, prostate and vesicles remain normal, and pressure applied directly to the testicle gives little or no pain. Tuberculous deposits tend to soften and to involve the scrotal coverings in spite of treatment. It is differentiated from haematocoele by the history of injury, or by the absence of the history or signs of syphilis. It can be
distinguished from *malignant tumours of the testis* by the history of syphilis, the tendency of syphilitic disease to be bilateral, the slow enlargement and positive serological tests. In malignant disease, the increase in the size of the testicle is more rapid, while the tumour often shows areas of varying consistency; the cord is often thickened in malignant or in tuberculous cases, but seldom in syphilitic ones.

**AFFECTIONS OF THE COVERINGS OF THE TESTIS CAUSING PAIN IN THE ORGAN**

The only common lesions of the coverings of the testis are hydrocoele and haematocoele; tumours of the testicular tunics are so rare as to render them surgical curiosities, and they rarely cause pain.

**Hydrocoele**

Hydrocoele may occur occasionally as an acute lesion accompanying an acute epididymitis or injury to the scrotum, or in the course of acute specific fevers such as mumps. Acute hydrocoele has been described in conjunction with acute lesions of other serous membranes, for example polyserositis. The more usual form of hydrocoele is the chronic variety, which may be due to some disease of the testis, but for which, in the majority of cases, no ascertainable cause can be found (primary or idiopathic hydrocoele).

A hydrocoele is usually painless but may cause some aching in the testicle, or a dragging sensation in the inguinal or iliac areas from the mechanical effect of its weight. It forms a swelling on one side of the scrotum, oval with a smooth uniform surface; it gives a distinct sense of fluctuation. The swelling is limited above from the cord or external abdominal ring and gives no sense of impulse on coughing; with a good light, it can be found in most cases to be translucent, the testicle occupying a posterior and low position in the swelling. The diagnosis of hydrocoele is usually easy, but difficulty may be experienced in long-standing cases in which the walls are much thickened. A hydrocoele must be diagnosed from: (i) a scrotal hernia; (ii) a haematocoele; (iii) a tumour; and (iv) a cyst of the epididymis:

- **Scrotal hernia.** An indirect inguinoscrotal hernia usually gives an impulse on coughing, can be reduced into the abdomen with a sudden slip or gurgle, and varies in size with the position of the patient. A hernia comes from above and descends into the scrotum, the swelling is not limited above, and the testis can be distinguished at the bottom of the scrotum. A hydrocoele is distinctly limited above so that the examining fingers can meet above it, it gives no impulse on coughing, it is translucent, and the spermatic cord can easily be distinguished (see Fig. T3a). The testis in a hydrocoele cannot usually be distinguished in the scrotum, as in a hernia. Difficulty may arise between the two conditions when the hydrocoele extends along the funicular process in the inguinal canal, and thus gives an impulse on coughing, or if the translucency is lost owing to the thickness of the walls of the sac. A scrotal hernia in an infant may be translucent.

- **Haematocoele** is distinguished from hydrocoele by the absence of translucency and the rapidity of the onset, usually after an injury or puncture (see also below).

- **Tumours of the testis.** A hydrocoele has a much slower rate of increase in size, has a smooth surface and uniform consistency, and is translucent.

- **Cyst of the epididymis** (see above).

In cases of doubt, ultrasonography of the swelling usually enables accurate anatomical delineation of the mass to be made, and distinguishes between a cystic and a solid swelling.

**Haematocoele**

Haematocoele may occur from puncture of a vein in the sac or of the testicle as the result of tapping a hydrocoele, or by the occurrence of bleeding into a hydrocoele. It may occur quite independently of a hydrocoele, usually after direct injury. As a rule, there is a rapid onset of swelling in the scrotum following the injury, with ecchymosis of the scrotal skin; the resulting tumour resembles a hydrocoele in its clinical symptoms, save that it is not translucent. In other cases, the swelling arises more slowly, when an oval swelling is present in one side of the scrotum covered by normal skin; the surface of the swelling is smooth, and gives a sense of fluctuation and elasticity. There is no translucency and, on tapping, dark blood-stained fluid is withdrawn.

The diagnosis in the less acute cases often presents a difficulty, especially with regard to *malignant disease of the testicle* (see above); this is particularly so when the haematoma is organized. It is distinguished from a hydrocoele by the absence of translucency, and from a hernia by the same points, except translucency, mentioned above in the diagnosis between hydrocoele and hernia.

**AFFECTIONS OF THE SPERMATIC CORD CAUSING TESTICULAR PAIN**

An inflammatory affection of the cord secondary to urethral infection is not uncommon. Tuberculous infection of the cord is practically never present without corresponding infection of the epididymis.
New growths of the cord, lipomas, sarcomas
(extremely rare) and hydrocoele of the cord cause
no pain in the testis. A varicocele, especially if large,
in a pendulous scrotum is a frequent cause of a dull,
aching pain in the testicle; it is nearly always left-
sided, although the reason for this is obscure. The
characteristic feel of the enlarged veins in the erect
position (like a ‘bag of worms’), and the slight impulse
and thrill on coughing, will readily point to the correct
diagnosis.

**UNDESCENDED TESTIS**

This condition should ideally be recognized and treated
in early childhood before the age of 2 years. It may
give rise to pain. A testis may be arrested in its descent
at the external abdominal ring (superficial inguinal
pouch) or in the inguinal canal, may remain inside the
abdomen, or may pass upwards and outwards from
the external abdominal ring into the superficial inguinal
pouch, where it can be felt readily. Occasionally, it
passes into the perineum after traversing the inguinal
canal, to the upper part of the thigh via the crural ring,
or to the root of the penis in front of the pubis. In one-
fifth of all cases, the undescended testes are bilateral.

An undescended or ectopic testicle may be involved
with the range of pathology that also affects the
normally placed organ, and thus give rise to pain.
However, in addition, owing to the effect of recurrent
muscular strains and the comparative immobility of the
organ, it is particularly liable to attacks of inflammation.

In the intra-abdominal position, it remains protected
from muscular injury, while ectopic testicles have a
greater range of mobility than has one that is retained
in the inguinal canal and are thus especially prone to
torsion. The inflammation of an undescended testicle
may be so acute as to lead to gangrene of the organ,
with or without torsion of the cord. The pain may be
complained of first when the testes begin to enlarge at
puberty, at which time an undescended right testicle
may produce symptoms that can be easily mistaken for
appendicitis.

The diagnosis of undescended testicle rests upon the
following points: the fact that one side of the scrotum
is empty, the outline and situation of the swelling in the
superficial inguinal region or elsewhere, the testicular
sensation upon pressure, and the recurrent attacks of
pain. An undescended testicle may give rise to acute
pain from inflammatory lesions or from acute torsion
of the organ, and may, if it is in the inguinal canal, give
rise to symptoms suggesting a strangulated hernia.

A partially descended testicle is often accompanied by
an inguinal hernia. The misplaced testis is especially
liable to result in malignant disease.

It should be remembered that an imperfectly
descended testis is a small, atrophic and poorly
developed organ, and the spermatogenesis from the
gland is usually absent.

**TESTICULAR PAIN FROM LESIONS OTHER THAN IN THE TESTICLE**

Complaint may be made of testicular pain when,
on clinical examination, the testis is found to be
normal. After an acute inflammation of the organ,
even when no palpable nodule remains, the resulting
inflammation may cause aching in the organ, especially
after sexual excitement or prolonged desire. Apart
from former testicular disease, referred pain may be
felt in the organ if a calculus is present in the pelvis
of the kidney or upper ureter, or from stimulation
of the peripheral nerves by secondary deposits in
the bodies of the lumbar vertebrae, pressure from an
extramedullary intraspinal tumour such as a
neurofibroma, meningioma or ependymoma, or
the pressure of an aneurysm in this situation. Pain
in the testis is occasionally present in appendicidal
inflammation when the appendix turns down into the
pelvis. Finally, when no organic cause of any sort is
present the condition is usually called neuralgia testis;
this is pain of an aching character that may occur in
patients of a neurotic tendency.

**TESTICULAR SWELLING**

Ben Challacombe

(See also TESTICULAR PAIN, p. 662.)

It is first essential to prove that the swelling is really
testicular and not an inguinoscrotal hernia. This is
done by grasping the root of the scrotum between the
thumb and index finger to determine whether any of
the swelling extends along the cord into the inguinal
region. True scrotal swellings may arise in: (i) skin; (ii)
the various connective tissue coverings of the testicle;
(iii) the tunica vaginalis; (iv) the testis; (v) the epididymis;
and (vi) the lower end of the spermatic cord.

**SWELLINGS AFFECTING THE SKIN**

The nature of these is usually obvious. The only
common ones are sebaceous cysts. Much less common
are soft sores, chancre, warts and epithelioma. The last-
named soon ulcerates, and it was once commonly seen
in sweeps or in those who worked in tar, tar products
or petroleum. It is now relatively rare.

**SWELLINGS OF THE VARIOUS CONNECTIVE TISSUE COVERINGS**

These are rare, but occasionally a fibrosarcoma may
occur. These swellings are movable upon the testicle.
and not attached to it. The symmetrical enlargement called elephantiasis scroti (Fig. T.2), due to *Wuchereria bancrofti*, is limited to filarial infection in the tropics, although sometimes a similar state of scrotal distension and overgrowth results in from lymphatic obstruction due to pelvic cellulitis, abdominal malignancy or congenital abnormality. The enlarged scrotum resulting from acute generalized oedema in acute or chronic renal disease, hepatic or cardiac failure or hypoalbuminaemia is seldom difficult to recognize; the penis and prepuce are generally distended by oedema at the same time as are the legs, loins, eyelids and other parts, and the diagnosis is confirmed by the albumin and tube-casts in the urine. Gross oedematous scrotal swelling also occurs with ascites or inferior vena caval thrombosis, or after inguinal/femoral lymph node dissections, and it may accompany the abdominal swelling of pellagra and infantile kwashiorkor.

**THE TUNICA VAGINALIS**

The tunica vaginalis may become distended with serous fluid, blood or pus; distension with fluid may be primary, the idiopathic vaginal hydrocele, or secondary to disease of the testis or epididymis. The common vaginal hydrocele (within the tunica vaginalis) usually arises slowly, although some directly follow injury. The patient is well, with no pain or urinary complaint, and merely complains of the lump or of the drag it causes. The swelling is large, heavy, ovoid, tense and elastic rather than fluctuating, although fluctuation can be proved if the swelling is fixed by an assistant or the patient; neither testis nor epididymis can be felt separately from the swelling. A hydrocele glows red when transilluminated, but it needs a dark room and a strong light (Fig. T.3): when transilluminated the testicular shadow will be noticed at one edge of the swelling, usually behind it. Aspiration is generally not recommended due to the risk of introducing infection, 

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**Figure T.2** Elephantiasis of the scrotum due to filariasis. [Dr C.J. Hackett, Wellcome Museum of Medical Science.]

**Figure T.3** (a) Left vaginal hydrocele; the fingers reach above it, thus excluding inguinoscrotal hernia. (b) The hydrocele transilluminates brilliantly.
but it produces a golden-or straw-coloured fluid. A secondary hydrocele follows disease (infection, tumour or trauma) to the testis or epididymis; the amount of fluid is usually small, and the swelling lax, so that the finger can be passed through it to touch the testis. The complaint is of the causative disease rather than of the hydrocele, which is usually discovered on examination or ultrasound assessment. A haematocele has the physical characters of a hydrocele except that it is not translucent, and it contains blood. A haematocele is due to injury, torsion or occasionally tumour of the testis, and ultrasound or tapping may establish the presence of blood. A pyocele is merely part of a suppurative process arising in the testis or the epididymis. The differential diagnosis of hydrocele is from translucent swellings in the epididymis and cord, cysts of the epididymis and encysted hydrocele. Ultrasonography has proved to be invaluable in the investigation of testicular masses, in particular in determining whether or not there is underlying testicular pathology in a patient with a hydrocele.

**Swellings of the Testis**

These usually affect either the body of the testis or the epididymis, and rarely the two together. The first group includes torsion, mumps, gumma and tumours; the second tuberculosis, gonorrhoea, bacterial (often *Escherichia coli*) infection and cysts. Determination of the anatomical site of the swelling will usually point towards its pathological nature.

**Swelling of the body of the testis**

*Torsion* is an acute condition accompanied by severe scrotal or abdominal pain and vomiting, and it occurs frequently in the undescended testis (see p. 666). Torsion of a fully descended testis, giving rise to a scrotal swelling, is seldom seen after adolescence; the local signs, in addition to the abdominal pain and vomiting, are moderate enlargement of the testicle, acute tenderness, the presence of a small haematocele, and the appearance after a few hours of oedema of the scrotal wall on the affected side. The testis may have a horizontal lie as opposed to the normal vertical position, when it is said to resemble a ‘bell-clapper’. Recurring or intermittent torsion of the testicle is not uncommon, and in these cases the signs and symptoms are less pronounced than in the acute variety, into which they eventually pass. The main points of distinction between the less acute enlargements of the corpus testis are described in Table T.1.

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**Table T.1 Swellings of the body of the testicle**

<table>
<thead>
<tr>
<th>Swelling</th>
<th>Mumps</th>
<th>Syphilis</th>
<th>Tumour</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Age</strong></td>
<td>Puberty or adolescence</td>
<td>Any age, but usually 18–30 years</td>
<td>Any age, more common after 20 years</td>
</tr>
<tr>
<td><strong>History and other symptoms</strong></td>
<td>Short history with pyrexia</td>
<td>Previous history of exposure to venereal disease; usually has had chancre and rash</td>
<td>Onset insidious</td>
</tr>
<tr>
<td></td>
<td>Previous contact with mumps</td>
<td>Gumma or tertiary rashes may be found elsewhere</td>
<td>History of months</td>
</tr>
<tr>
<td></td>
<td>Parotids enlarged</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Scrotum</strong></td>
<td>Normal or red and hot</td>
<td>Normal or adherent in front Later, ulcer with sharp edges and slough at base, or hernia testis</td>
<td>Normal or merely stretched until growth is size of tennis ball, when it may be invaded</td>
</tr>
<tr>
<td><strong>Testis</strong></td>
<td>Moderately enlarged, shape normal</td>
<td>Enlarged; up to two or three times normal May be nodular</td>
<td>Increases steadily and may reach diameter of 10–13 cm Initially smooth, later nodular</td>
</tr>
<tr>
<td><strong>Sensation</strong></td>
<td>Tender and painful Testicular sensation present</td>
<td>Not tender or painful Testicular sensation lost</td>
<td>Painful, but not tender</td>
</tr>
<tr>
<td><strong>Tunica vaginalis</strong></td>
<td>Slight hydrocele in most</td>
<td>Hydrocele in 60%</td>
<td>Hydrocele in early stages; later haematocele</td>
</tr>
<tr>
<td><strong>Epididymis</strong></td>
<td>Unaltered</td>
<td>Usually unaltered</td>
<td>Flattened</td>
</tr>
<tr>
<td><strong>Cord</strong></td>
<td>May be tender</td>
<td>Normal</td>
<td>Usually normal but may have nodules of growth in lymphatics</td>
</tr>
<tr>
<td><strong>Lymph nodes</strong></td>
<td>Not characteristically enlarged</td>
<td>Not characteristically enlarged</td>
<td>Drainage to para-aortic nodes at kidney level. These may form a very large mass Eventually, left supraclavicular nodes involved. Inguinal nodes not enlarged unless scrotal skin is invaded</td>
</tr>
</tbody>
</table>
Mumps orchitis usually occurs within a week of the parotid swelling and results in an acutely tender enlargement of one or both testes. It especially complicates the acute parotitis of adolescents or adults after 3–4 days. The affected testis often undergoes atrophy. If the condition is bilateral (10–30 per cent), it may result in infertility through tubular atrophy.

It is classically difficult to distinguish syphilitic enlargement of the testicles from that due to malignant tumour, but a course of antisyphilitic therapy and the serological reactions may resolve the uncertainty. Gumma of the testis, once common, is now a clinical rarity in Western communities (see also TESTICULAR PAIN, p. 662), therefore a solid mass in the body of the testis is highly suspicious of a neoplasm.

**Tumours of the testis**

Primary testicular tumours are the most common solid malignancy in men aged 25–45 years. Nearly all tumours of the testis are highly malignant. Any solid mass within the body of the testis should be considered a tumour until proved otherwise. Primary tumours are classically painless, but up to 5 per cent occur with acute scrotal pain or present after minor trauma. They fall into two main pathological varieties: seminoma (48 per cent) and non-seminomatous germ cell tumours (NSGCTs) (42 per cent), the latter including the subgroups of teratoma, yolk sac tumour, chorionic carcinoma and mixed non-seminomatous tumours. There is a smaller group of non-germ cell (sex-cord stromal tumours) neoplasms including Leydig cell, Sertoli cell and mixed or unclassified varieties of tumour. Rarely a Leydig cell tumour occurs in a child and produces sexual precocity.

Ultrasoundography of the testis can be very useful in localizing the exact site of the mass, and in differentiating between a solid and a cystic mass. If any doubt exists, it is advisable to excise the testis through an inguinal incision, first dividing the cord at the internal ring (Fig. T.4). Incision into a testis containing a malignant tumour is almost invariably followed by rapid recurrence. Tumour markers should be measured in the blood; human chorionic gonadotrophin (hCG), alphafetoprotein and lactate dehydrogenase may be elevated.

The seminoma (Fig. T.5) is a soft vascular solid growth composed of large spheroidal cells derived from the germinal epithelium of the seminiferous tubules. It is pale and homogeneous on sectioning. It commonly occurs in the fourth decade and is less malignant but more radio-sensitive than teratoma. It tends to retain the shape of the testis as it enlarges.

**Figure T.4** Seminoma of the testis.

**Figure T.5** Cross-section of a radical inguinal orchidectomy specimen showing two non-seminomatous tumours.

The teratomas (Fig. T.6) and other NSGCTs are solid or multinodular cystic growths in which one or other of the germinal layers may preponderate. NSGCTs are often heterogeneous and sometimes contain hair or teeth. Metastasis to the testis is rare, while lymphomas of the testis are the most common type of testicular tumour in men over 50 years old and are often bilateral. Most cases of teratoma occur at the age of 25–30; they give a short history, are more resistant to radiotherapy treatment, and show early metastases. There may be gynaecomastia, and chorionic hormone
(beta-hCG) may be present in the urine. In seminoma, the pituitary gonadotrophic hormone is sometimes found in the urine. The undescended testis is more prone to developing malignant disease than is the normally placed one. The testis that has been initially undescended and subsequently brought down into the scrotum maintains this higher tendency to malignant change, estimated at about 10 times that of the normal organ.

A testis containing a malignant growth enlarges slowly or rapidly but, as pain is at first absent, there may be nothing to arouse the patient’s suspicions. A rapidly growing malignant tumour of the testis may be so soft as to appear to be a fluid collection in the tunica vaginalis. Generally, however, although a growth may be accompanied by a small amount of fluid in the tunica vaginalis, the more solid mass can be felt through the fluid on careful examination. The epididymis may become incorporated in the growth so that it cannot be distinguished, and the tissues of the cord become thickened. The coverings of the testis become stretched over the tumour; the mass does not become adherent to the scrotal skin until late in the disease.

Clinically it is impossible to distinguish between a teratoma (NSGCT) and a seminoma. In both types, the para-aortic lymph nodes become enlarged, and they may be felt in a thin subject to one or other side of the epigastria area; pain due to the pressure of these nodes upon nerve structures may become marked. The inguinal nodes are usually not enlarged unless the scrotal skin is affected; retrograde spread may then occur to the iliac nodes, which may be felt at the brim of the pelvis. In advanced cases, the left supraclavicular lymph nodes are involved and become palpably enlarged. Mediastinal or pulmonary metastases are frequent, the latter giving the characteristic radiological appearance of ‘cannon-ball’ secondaries.

**THE EPIDIDYMIS**

The epididymis may become enlarged as the result of inflammation (usually infection), tumour or cystic degeneration. This usually occurs acutely, and the swelling may last 4–6 weeks. Primary tumours of the epididymis are excessively rare, and need not give rise to much concern in differential diagnosis.

Inflammatory swellings are characterized by being elongated in a vertical direction, by their relation to the testis, which they overlap at its posterior border and its upper and lower poles, and by being flattened from side to side. There may be an associated small, lax secondary hydrocele. Inflammatory swellings may be due to bacteria (commonly *Escherichia coli* but also other coliforms and Gram-negative organisms: *Mycobacterium tuberculosis, Chlamydia trachomatis, and Neisseria gonorrhoeae* (gonorrhoea)). Schistosomiasis may cause chronic painless epididymitis, while the antiarrhythmic drug amiodarone may cause unilateral or bilateral inflammation. The main points of the distinction are shown in Table T.2.

In tuberculosis, the epididymis feels like a beaded cord, and there may be thickening and beading of the vas deferens. If any doubt exists between the diagnosis of acute epididymitis and testicular torsion, an urgent surgical exploration is recommended. The treatment of epididymitis consists of culturing the urine and blood (if the patient is pyrexial), and prescribing suitable...
antibiotics (doxycycline or tetracycline for *Chlamydia trachomatis*, or ciprofloxacin/norfloxacin for *Neisseria gonorrhoeae*). Acute epididymitis can be complicated by testicular abscess, testicular infarction, chronic epididymitis and pain. Support of the testicle in a scrotal support or suspensory bandage will aid the symptoms.

Cysts of the epididymis may be solitary or multiple, and may be bilateral. A cyst of the epididymis is placed above and behind the testis, from which it is distinct. Although attached to the epididymis, it is rounded but, being thin-walled, it does not feel as tense as a hydrocele; it tends to have several rounded projections rather than a simple surface. Similarly to vaginal hydroceles, aspiration is not recommended due to the risk of infection, but any fluid withdrawn is milky or opalescent and shows numerous cells under the microscope, some of which may be spermatozoa. *Multiple cysts* occur in men past middle age. They are painless and increase in size very slowly. These swellings are strikingly translucent.

**SWELLINGS OF THE LOWER END OF THE CORD**

The most important swelling of the lower part of the spermatic cord is a *varicocele*. It may be mistaken for an inguinal hernia but has a characteristic feel like a ‘bag of worms’. It should be examined with the patient initially standing and subsequently lying. The swelling will reappear after it has been completely reduced by elevation of the scrotum and a finger released from firmly pressing on the external abdominal ring. The varicocele usually has a cough impulse and increases with a Valsalva manoeuvre. Varicoceles are far more common on the left than the right and, if of recent onset, may indicate obstruction to the drainage of the testicular vein by a renal tumour.

**TETANY**

Paul Carroll

*(See also CRAMPS, p. 117; TICS p. 682; VERTIGO, p. 736.)*

Tetany is a condition in which abnormal muscle cramps are caused by increased neuromuscular excitability induced by a fall in the concentration of ionized calcium. Important precipitating factors include a low-calcium diet, pregnancy and lactation. Early symptoms of acute tetany include paraesthesiae and numbness of the extremities and around the mouth. Then muscle cramps may occur, particularly in the extremities. The wrist and elbow may become flexed, the fingers flexed at the metacarpophalangeal joint, but extended at the terminal interphalangeal joint, and the fingers all pressed together, with the thumb adducted into the palm of the hand so that the whole hand forms the
‘main d’accoucheur’ (obstetrician’s hand). When the legs are affected, the knees and ankles are extended, the foot is arched and inverted, and the toes are flexed and pressed together. The cramps may be painful and last from several minutes to a couple of hours. In severe cases, the face and neck muscles may be involved, and laryngospasm may cause respiratory obstruction and lead to loss of consciousness and death. Generalized epileptic convulsions may occur, especially in children. Hypocalcaemia may also lead to prolongation of the Q-T interval on the electrocardiogram and predispose to ventricular tachyarrhythmias.

Latent tetany may be revealed by special signs:

- **Chvostek’s sign.** Twitching of the muscles of the upper lip may be elicited by tapping on the facial nerve in front of the ear. However, note that this sign can be present in some people with normal calcium levels.
- **Trousseau’s sign.** Spasm of the fingers and hand to form the ‘main d’accoucheur’ is induced by obstruction of the brachial artery for up to 3 minutes by a sphygmonanometer cuff.

Additional features, which may be present in patients with chronic tetany, include:

- Ectodermal changes (particularly in idiopathic hypoparathyroidism). The skin is dry and scaly, the tongue atrophic, the hair sparse and brittle, and the nails ridged; cataracts may be present.
- Central nervous system changes: general fatigue, irritability, anxiety, depression, paranoia and epilepsy. Papilloedema is sometimes found.

**CAUSES OF TETANY**

The causes of reduced serum calcium (hypocalcaemia) are listed in Box T.4.

**Box T.4 Causes of hypocalcaemia**

**Most common**
- Chronic renal failure
- Postoperative (thyroidectomy, parathyroidectomy)
- Osteomalacia/rickets

**Less common**
- Acute pancreatitis
- Alkalosis
- Magnesium deficiency
- Massive blood transfusion
- Drug therapy: bisphosphonates, calcitonin, phosphates

**Rare**
- Vitamin D resistance
- Pseudohypoparathyroidism
- Idiopathic hypoparathyroidism: autoimmune/congenital (DiGeorge syndrome)

**Renal failure.** In acute renal failure, hypocalcaemia may occur secondary to phosphate retention and the formation of insoluble calcium compounds. In chronic renal failure, calcium levels may fall due to a fall in the renal hydroxylation of vitamin D.

**Vitamin D deficiency.** Skin irradiation is the most important source of vitamin D, blood levels of which rises in the summer and fall in the winter. Elderly people and certain ethnic groups who do not venture outside very much are most at risk of vitamin D deficiency. It is difficult to provide enough vitamin D from even a normal diet. Vitamin D is found in wheatgerm, eggs and fish. Margarine and milk are fortified with vitamin D in some countries. Malabsorption syndrome of all types may cause vitamin D deficiency. Since vitamin D must be hydroxylated in the liver and kidney in order to create its most potent form, diseases of these organs may lead to hypocalcaemia. The problem of vitamin D deficiency is always exacerbated by prolonged lactation.

**Acute pancreatitis.** Hypocalcaemia may occur early on in acute pancreatitis. The mechanism is not clearly understood, but part of it may be due to the formation of calcium soaps.

**Magnesium deficiency.** This leads to reduced secretion of parathyroid hormone (PTH) and to peripheral resistance to the actions of PTH. It occurs in: (i) patients on prolonged intravenous therapy who are not receiving magnesium supplements; (ii) severe diarrhoea; (iii) malabsorption syndrome; (iv) small-bowel resection or bypass; (v) chronic alcoholism; (vi) the diuretic phase after recovery from renal tubular necrosis; and (vii) hyperaldosteronism.

**Alkalosis.** This condition, from any cause, may be associated with hypokalaemia and can lead to a decrease in the ionized fraction of calcium and cause tetany. The causes of alkalosis are: (i) the excessive ingestion of alkali; (ii) frequent vomiting; (iii) hyperventilation, especially hysterical; and (iv) hyperaldosteronism, where magnesium deficiency also plays an accessory role.

**Pseudohypoparathyroidism.** This is a rare disorder in which the hypocalcaemia is due to end-organ resistance to the actions of PTH, the concentration of which is raised. The patients have certain phenotypic characteristics (see also STATURE, SHORT, p. 639).
Disorders of thinking are commonly considered under two separate headings:

- **Disorders of the content of thinking**: these include adherence to delusional beliefs, as found in psychotic individuals, and the presence of unwelcome, intrusive obsessional thoughts found in patients with obsessional neurosis. These are discussed in other sections (DELIUSIONS, p. 131; OBSESSIONS, p. 470).

- **Disorders of the thinking process**, including abnormalities of the flow and the logical form of thoughts; this category will be considered more fully here.

A person's thought processes can only be inferred from their words and actions. We may consider that a person's thoughts are racing when they have pressure of speech or flight of ideas, as in mania. Here, there are logical connections between the ideas expressed, but the subject's thinking is so speeded up that it tends to jump from topic to topic, often in response to environmental distractions. The speaker is difficult to interrupt, loud, emphatic and may use clang associations, when the choice of words is governed by their sound rather than by a logical relationship. Sentences may be linked by rhyming and punning, often with amusing consequences.

The above description is typical of mania, but lesser degrees of increased flow of thinking may be found in acute reactions to stress, major depressive illness with marked features of anxiety, and occasionally schizophrenia and organic mental disorders. A reduced flow of speech may reflect poverty of thought, when the subject makes no spontaneous conversation and responds to questions with only brief unelaborated replies in a flat monotonous tone. In its most severe form, it may lead to muteness, and this occurs frequently in major depressive episodes, schizophrenia and the dementias.

Disorders in the form of thinking (formal thought disorder) are of enormous importance in the diagnosis of schizophrenia, but similar features may also occur in normal subjects who are tense or fatigued, and in patients with organic brain diseases. An important symptom in schizophrenia, and one that can often be observed in patients with long-standing chronic disability, is loosening of association of thoughts, leading to incoherence of speech. These characteristic changes may be accompanied by other so-called negative symptoms of schizophrenia, including a degree of apathy with diminished drive and volition, which may be the principal feature of the disease after the more florid symptoms of delusions and hallucinations have resolved in response to antipsychotic medication. Loosening of association describes a thought disorder characterized by speech in which ideas shift from one subject to another completely unrelated topic without the speaker being aware that the topics are disconnected. Unlike flight of ideas, the flow of talk may be normal or even reduced. Derailment is a description sometimes used for such idiosyncratic moves from one frame of reference to another; if severe, speech becomes word salad and totally incoherent. In very severe schizophrenic thought disorder, the speech has some of the characteristics of a fluent aphasia (schizophrenia), becoming a series of totally incomprehensible sentences often admixed with neologisms (invented words or word combinations). Thought-blocking is expressed as an interruption in the normal flow of speech, usually in a patient with schizophrenia. The subject, moreover, is aware and will spontaneously describe how their train of thought was suddenly stopped and they had no control over it. The symptom is quite different from mere absentmindedness, and sometimes the patient may elaborate the paranoid delusion that an external force is responsible for taking their thoughts away. Concrete thinking is again a term frequently applied to schizophrenic patients, but it is also observed in people with a moderate learning disability, and among brain-damaged and autistic individuals. It describes a difficulty in handling abstract and symbolic language leading, for example, to very literal interpretations of proverbs. When mild, this feature has no diagnostic value as it reflects cultural, educational and personality factors. In perseveration, found in organic brain disorders and in schizophrenia, the patient continues to hold ideas or tasks after they have ceased to be appropriate, so that the same word or idea will crop up in the patient's speech many times in a few sentences. It may be quite marked, for example, 'I think I'll put on my hat, hat, hat, hat', when the term 'logo clonus' or 'syllable repetition' would apply. Such problems switching paradigm, the so called 'executive function', point to dysfunction of the frontal lobes of the brain.

Obsessional and anxious people may betray aspects of their thinking by circumstantiality. Here, speech may be produced in a slow stream but with a great excess of unnecessary detail that can entirely obscure the main answer to the question. Circumstantiality, however, is commonly found in those with no mental illness.
mediastinitis account for some cases, the latter
Large retrosternal goitres or chronic fibrous
innominate artery are now rare.
Aneurysms of the ascending aorta and
metastatic carcinomas may also give rise to SVC
thymomas, germ-cell tumours and, to a lesser extent,
and mediastinal infiltration. Lymphomas, malignant
lobe and causing paratracheal gland enlargement
Malignant disease accounts for most cases, especially
dysphagia, headache, stupor, seizures and syncope.
1757 in a patient whose SVC was obstructed by a
condition was first described by William Hunter in
obstruction of the superior vena cava (SVC). The
This may be caused by either partial or complete
azygos veins.
Venous blood from the posterior chest wall muscles,
skin and vertebral venousplexuses drains into
11 intercostal veins. On the right side, the
second, third and often fourth posterior intercostal
veins unite to form the right superior intercostal vein,
which then drains directly into the azygos vein. The
lower posterior intercostal veins individually drain into
the azygos vein. On the left side, the second, third and
sometimes fourth posterior intercostals veins unite to
form the left superior intercostal vein, to open directly
into the left innominate vein. The lower left posterior
intercostal veins drain individually into the hemiazygos
vein, to then cross the midline behind the mediastinal
structures to the azygos vein, and on to the superior
vena cava just above the pericardium. The superior
vena cava arises from the innominate, jugular and
azygos veins.
The thoracic wall veins may become distended in a
number of conditions, usually associated with partial or
complete obstruction of blood flow in the innominate
or vena cava to cause a rise in pressure in the azygos,
hemiazygos or jugular venous systems.
SUPERIOR VENA CAVAL SYNDROME
This may be caused by either partial or complete
obstruction of the superior vena cava (SVC). The
condition was first described by William Hunter in
1757 in a patient whose SVC was obstructed by a
sYPHILITIC AORTIC ANEURYSM. The signs and symptoms
can be subtle and may evolve slowly over a few
weeks. Characteristic signs include cyanosis, oedema,
venous engorgement of the head, neck and arms,
chest and upper abdomen, brawny non-pitting
oedema of the neck and dysphonia due to laryngeal
oedema. Symptoms, which frequently worsen when
the patient lies down or leans forward, include facial
congestion and swelling, breathlessness, cough,
dysphagia, headache, stupor, seizures and syncope.
Malignant disease accounts for most cases, especially
from small-cell carcinoma involving the right upper
lobe and causing paratracheal gland enlargement
and mediastinal infiltration. Lymphomas, malignant
thymomas, germ-cell tumours and, to a lesser extent,
metastatic carcinomas may also give rise to SVC
obstruction. Benign lesions account for about 5 per
cent of cases. Aneurysms of the ascending aorta and
innominate artery are now rare.
Large retrosternal goitres or chronic fibrous
mediastinitis account for some cases, the latter
being secondary to tuberculosis, histoplasmosis,
coccidioidomycosis, blastomycosis or filarial
mediastinal lymphadenitis.
In a proportion of patients, the cause of mediastinal
fibrosis remains obscure. Iatrogenic causes of
SVC obstruction include thrombosis following the
introduction of subclavian lines for venous access
or the insertion of temporary pacemaking wires.
Superior vena caval obstruction is seldom a medical
emergency, but it does warrant prompt investigation
and treatment. Whenever possible, a histological
diagnosis should be obtained, either by bronchoscopy,
mediastinoscopy or thoracotomy, to facilitate effective
treatment for lymphoma, small-cell carcinoma, other
tumours or non-malignant conditions.
Lesser degrees of localized chest wall venous distension
may occur secondary to axillary vein thrombosis.
This is not an uncommon condition and may occur
after vigorous use of the arm, allowing the vein to be
compressed and damaged between the clavicle and
first rib. Painful congestion and oedema of the affected
arm follow, with collateral vein distension on the upper
chest. The condition usually settles over 3 months.
Thrombophlebitis of the superficial veins of the
breast and anterior chest wall (Mondor’s disease) are
also encountered. This condition may also affect the
arm. The essential characteristic physical sign is an
indurated subcutaneous thrombophlebitic cord about
3 mm in diameter. The cause is uncertain, and the
condition gradually subsides over a few months.

THRILLS

Alex West

THORACIC WALL VEINS

Venous blood from the posterior chest wall muscles,
skin and vertebral venousplexuses drains into
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indurated subcutaneous thrombophlebitic cord about
3 mm in diameter. The cause is uncertain, and the
condition gradually subsides over a few months.

THRILLS

Gerry Can-White

(ASee also HEART, MURMURS IN, p. 269.)

A thrill is a palpable vibration of vascular origin.
It indicates vascular turbulence close to the site
where it is felt. Thrills can be divided into: (i) those
of cardiac origin, which can be regarded virtually as
palpable heart murmurs; and (ii) those which arise in
the extracardiac vessels. Cardiac thrills are discussed,
along with heart murmurs, on page 269. Extracardiac
thrills are usually accompanied by an audible
component, termed a ‘bruit’.

Carotid thrills usually arise from turbulence generated
at the aortic valve rather than from a local carotid
stenosis. This is perhaps because turbulence in
the relatively small carotid vessel generates a
high-frequency sound that is more easily heard
than felt. A slow-rising carotid pulse with a thrill is
generally pathognomonic of aortic stenosis, and this

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is confirmed by hearing an aortic ejection systolic murmur and demonstrating aortic valve disease by electrocardiography. It must be remembered that intracardiac and carotid lesions may co-exist.

Subclavian thrills may be generated locally or arise from the aortic valve. A difference in timing of the pulse at the two wrists, and unequal blood pressures in the arms, should be sought. A subclavian aneurysm may cause a thrill, but these are rare.

Peri-scapular thrills are a feature of aortic coarctation and are due to dilatation and increased flow in the peri-scapular vessels, as these form part of an anastomotic circulation around the aortic obstruction. There is often an associated bruit, together with upper segment hypertension and delayed or absent femoral pulses.

Cimino–Brescia arteriovenous fistulae created surgically in the vessels of the forearm (less commonly the leg), to allow easy venous access for haemodialysis in patients with renal failure, are probably the most common cause of peripheral vascular thrill in modern practice. The presence of a thrill indicates that the anastomosis remains functional. Sometimes these fistulae enlarge excessively and have adverse haemodynamic effects. Congenital arteriovenous fistulae are much less common. In children, they may cause excessive lengthening of a limb. There is usually a conspicuous dilatation of superficial vessels.

Abdominal thrills are uncommon. An arteriovenous malformation in the liver, or a very vascular tumour such as an angiosarcoma, may cause a palpable thrill usually accompanied by a bruit.

Femoral artery thrills are usually the result of iliac atheroma and are accompanied by a bruit and evidence of peripheral vascular disease elsewhere. Very occasionally, they may be due to a femoral arteriovenous fistula, which is a rare complication of cardiac catheterization by the femoral route.

**THROAT, SORE**

Michael Gleeson

Sore throats affect everyone from time to time. Most are part of a viral upper respiratory infection. Others do not progress in that way, and the infection remains restricted to the pharynx. Although a large number of these infections have a viral cause, other sore throats are caused by bacterial infection, and an increasing number are fungal. The development of a persistent sore throat in a middle-aged or elderly smoker should raise concern, as it might be the first symptom produced by an upper aerodigestive tract neoplasm. A past history of excessive spirit consumption should heighten this concern and prompt a thorough clinical examination of the oral cavity, oropharynx, hypopharynx, larynx and neck. The pain experienced on swallowing in this group of patients often radiates to the ear.

The causes of sore throat are listed in Box T.5. Influenza, herpes simplex, adenosivirus, rhinovirus, Coxsackie and Epstein–Barr viruses cause viral pharyngitis. Patients usually complain of severe pain with, perversely, very little to find on clinical examination. There will probably be redness and oedema of the pharynx, especially along the pillars of fauces, the tonsils and the posterior pharyngeal wall. Lymphoid aggregates in the pharyngeal mucosa swell to produce a nodular or granular appearance on the posterior pharyngeal wall. Symptomatic relief with analgesics and mouthwashes is all that can be offered, other than sympathy. The condition resolves spontaneously in about 2–3 days.

Apart from pharyngitis, acute tonsillitis is probably the most common cause of a sore throat. This condition mainly affects the young, with peak incidences between 4 and 6 years of age, and again later in adolescence. Acute tonsillitis presents with malaise, pain and dysphagia. Patients are pyrexial and have enlarged tonsils with pus exuding from the crypts, especially along the pillars of fauces, the tonsils and the posterior pharyngeal wall. There will probably be redness and oedema of the pharynx, especially along the pillars of fauces, the tonsils and the posterior pharyngeal wall. Lymphoid aggregates in the pharyngeal mucosa swell to produce a nodular or granular appearance on the posterior pharyngeal wall. Symptomatic relief with analgesics and mouthwashes is all that can be offered, other than sympathy. The condition resolves spontaneously in about 2–3 days. Complete recovery takes 5–7 days, considerably longer than recovery from a viral pharyngitis. However, some do

<table>
<thead>
<tr>
<th>Box T.5 Causes of sore throat</th>
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<tbody>
<tr>
<td><strong>Tonsillitis</strong></td>
</tr>
<tr>
<td>• Bacterial: streptococcal</td>
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<tr>
<td>• Viral: infectious mononucleosis</td>
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<tr>
<td><strong>Pharyngitis</strong></td>
</tr>
<tr>
<td>• Viral: rhinovirus, Coxsackie, Epstein–Barr virus, adenosivirus, herpes, HIV</td>
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<tr>
<td>• Fungal: Candida, phycomycetes, blastomycetes</td>
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<tr>
<td><strong>Mucocutaneous disorders</strong></td>
</tr>
<tr>
<td>• Aphthae, lichen planus, pemphigus, pemphigoid, Behçets disease</td>
</tr>
<tr>
<td><strong>Neuralgias</strong></td>
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<tr>
<td>• Glossopharyngeal</td>
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<tr>
<td>• Eagle’s syndrome</td>
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<tr>
<td>• First bite pain</td>
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<tr>
<td><strong>Neoplasia</strong></td>
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<tr>
<td>• Carcinoma of the tongue, palate, pharynx and hypopharynx</td>
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<tr>
<td><strong>Miscellaneous</strong></td>
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<tr>
<td>• Thyroiditis</td>
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<tr>
<td>• Gastro-oesophageal reflux disease</td>
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<td>• Angina</td>
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</tbody>
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not respond to treatment, or present late when the infection has spread beyond the tonsillar capsule and pus has collected in the peritonsillar space – a *quinsy*. These patients are extremely toxic, and have marked trismus and a very ‘plummy-sounding’ voice. Only a limited view of the oropharynx is possible in these unfortunate patients, but it is usually sufficient to appreciate the swelling and faucial oedema of one tonsil, together with displacement of the uvula to the other side (Fig. T.7). Apart from being very unpleasant, this can develop into an extremely serious condition if the infection continues to spread and involve the parapharyngeal space; this is termed *parapharyngeal space abscess*.

*Infectious mononucleosis* (*glandular fever*) is caused by infection with the Epstein–Barr virus. In this condition, which normally affects adolescents, there is malaise, pyrexia, tonsillitis and a marked cervical lymphadenopathy. The tonsils are covered with a thick, white plaque, and there may be petechial haemorrhages at the junction of the hard and soft palate (Fig. T.8). In some, the tonsillar enlargement can be so severe that the airway becomes compromised and the patient has stertor. Some patients will also have hepatosplenomegaly.

*Fungal pharyngitis* caused by *Candida albicans* is increasingly being seen. It is associated with the use of inhaled steroids (Fig. T.9), and, in an otherwise fit young man, may be one of the presenting features of HIV infection. Other immunocompromised patients are also susceptible to this form of infection. Less common is infection with the phycomycetes and *Blastomyces* fungi, and these are usually confined to selected geographical areas.

*Aphthous ulceration, Behçet’s syndrome, pemphigus, pemphigoid and erosive lichen planus* can all affect the mucosa of the pharynx. These conditions give rise to painful superficial ulceration that often also involves the mucosa of the oral cavity. In the case of pemphigus and pemphigoid, ulceration is preceded by a bullous lesion. The recurrent nature of these lesions and other cutaneous lesions suggests the diagnosis in most cases. Patients with major aphthae always have relatively few ulcers, and in some cases they may be solitary. Healing can take a number of weeks in these patients, and this may cause concern that the diagnosis might be incorrect. If in doubt, a biopsy is prudent (Fig. T.10).

A number of conditions can give rise to neuralgic pain in the throat that is either spontaneous or precipitated by swallowing. *Glossopharyngeal neuralgia* is the immediate counterpart of trigeminal neuralgia. Patients with this condition experience paroxysms of...
excruciating pain, sometimes spontaneous, but at other times triggered by eating or swallowing. In Eagle’s syndrome, a similar type of pain is also experienced when swallowing. It is caused by friction between the glossopharyngeal nerve and an elongated styloid process. First bite pain is again similar in character and affects those patients who have undergone surgery on carotid body tumours or the cervical sympathetic chain. Their pain is triggered by chewing food or swallowing. Characteristically, it only affects these patients during the first meal of the day, but in some it is more prolonged. Acute or subacute thyroiditis can cause pain in the neck or throat. In the acute phase of the disorder, the patient has no difficulty in localizing the problem to the neck. Patients with subacute thyroiditis, however, frequently complain of a persistent soreness in the throat. Discomfort is constant and aggravated by swallowing; it is associated with the sensation of a lump in the throat. Patients are intolerant of constriction of the neck, for example by shirt collars. Reflux oesophagitis usually causes vague symptoms of soreness in the throat, or a sensation of a lump in the throat caused by cricopharyngeal spasm. Patients may have chronic hoarseness or a constant feeling of wanting to clear their throat. The pain may be aggravated after meals or at night, when patients are recumbent and subject to free reflux.

Without doubt, the most important issue in middle-aged and elderly patients with a sore throat is to exclude a carcinoma. This almost always affects those who have smoked heavily for a prolonged period, many of whom will also have consumed large amounts of spirits. Most of these tumours can only be seen on mirror examination of the pharynx or with endoscopes (Fig. T.11). The pain is progressive, often radiates to the ear, and may become or be associated with a husky voice, haemoptysis, cervical adenopathy and weight loss. Tumours of the oropharynx are not restricted to the middle-aged smoker and drinker. Nowadays more and more HPV positive oropharyngeal tumours are presenting in a much younger population who have not necessarily smoked or been heavy drinkers. Likewise patients with AIDS may present with Kaposi sarcoma in their throats (Fig. T.12).

When the diagnosis of sore throat is not immediately obvious on physical examination, cultures for bacteria and viruses, haematological studies, lateral X-rays of the neck, barium studies of the pharynx, oesophagus and stomach, and computed tomography scanning of the neck and examination under anaesthetic may be required. If pain in the throat is initiated or accentuated by exertion, a thorough cardiac assessment is also wise, as angina can be experienced in this way. As stated previously, frequent recurrent infections of the pharynx...
THYROID ENLARGEMENT

may be an expression of immunodeficiency, as seen in the DiGeorge syndrome or AIDS. It is also seen in subclass deficiencies of immunoglobulins.

THYROID ENLARGEMENT

Harold Ellis

(See also NECK, SWELLING OF, p. 454.)

An enlarged thyroid gland gives rise to a swelling in the front of the neck, medial and deep to the sternocleidomastoid muscles and medial to the carotid vessels, which, if the swelling is large enough, are displaced laterally and backwards. The gland is connected intimately with the larynx so that it rises and falls with the larynx and trachea during deglutition. This sign alone is generally sufficient to establish the diagnosis of an enlarged thyroid gland. The only other lump in the neck that moves on swallowing is a thyroglossal cyst, which characteristically, and in addition, moves upwards when the patient protrudes the tongue. This is because of the attachment of the cyst by a fibrous strand extending to the foramen caecum of the tongue (see Fig. N.20).

Inspection with the patient at rest and on swallowing may alone be enough to render a diagnosis of thyroid swelling extremely likely. Palpation will confirm this, and is usually best performed while standing behind the patient with the patient’s neck flexed and relaxed. The lateral lobes are palpated with the appropriate sternocleidomastoid muscle relaxed. If the enlargement is only slight, help may be obtained by displacing the trachea towards the side being examined, when it is possible to introduce the fingers under the relaxed sternocleidomastoid to feel the posterior border of the lobe. The trachea and larynx may of course already be the subject of pathological displacement by pressure of the enlarged gland, and this should be determined at the time of palpation. The larynx should also be examined with a mirror for paralysis or asymmetry of the vocal cords. Vocal cord paresis will usually be accompanied by alteration in the voice and, if both cords are affected, possibly with dyspnoea and stridor as well. The possibility of pressure effects always requires investigation, and these may be enumerated as follows:

- Pressure on the trachea causing deviation or compression or both, with varying degrees of dyspnoea and stridor
- Pressure on the oesophagus causing dysphagia
- Pressure on nerves, usually the recurrent laryngeal nerves, producing various forms of vocal cord palsy with or without alteration in the voice, dyspnoea, stridor and a ‘brassy’ cough. The cervical sympathetic is occasionally involved, as shown by contracted pupil and ptosis (Horner’s syndrome). Such nerve palsies are almost invariably associated with invasive tumours of the thyroid gland
- Pressure on veins, giving rise to engorgement and the setting up of anastomotic channels, as a result of superior mediastinal obstruction from a large retrosternal extension of the gland
- Acute pressure symptoms may arise, or those already present may become acutely aggravated, by haemorrhage into a thyroid cyst

Retrosternal prolongation of the thyroid should not be forgotten. It can be recognized by a dullness on percussion over the manubrium, but this sign is unreliable. When the patient is asked to swallow or cough, it is sometimes possible to feel the lower limit of the gland as it rises; at the end of deglutition, it slips back behind the sternum (‘plunging goitre’). The thyroid in the neck may occasionally appear of normal size in the presence of a retrosternal enlargement, and in a few rare cases, the whole gland lies behind the sternum. Pressure symptoms are liable to be great when part or the whole of the gland is in this position, and sometimes the result of pressure on the great veins is seen in the presence of dilated anastomotic skin veins over the upper anterior part of the thorax. Radiographic examination is a most useful adjunct in the diagnosis of thyroid enlargement, showing both the presence of retrosternal prolongation and tracheal displacement and compression. Other aids may be useful with individual cases.

VARIETIES OF ENLARGEMENT AND THEIR DIFFERENTIAL DIAGNOSIS

It should be noted that the term ‘goitre’ is in common use; this simply means an enlargement of the thyroid, from whatever cause.

- Physiological enlargement: occurs at puberty and during menstruation and pregnancy and is usually symptomless
- Inflammatory enlargement:
  - Acute: in acute thyroiditis, symptoms include the usual signs of acute inflammation; this condition is rare
  - Chronic: tuberculosis, syphilis, Riedel’s disease; all of these conditions are rare
- Simple goitre (endemic and sporadic). Parenchymatous goitre, colloid goitre, nodular goitre, solitary (fetal) adenoma
- Hyperthyroid (thyrotoxic) goitre:
  - Primary hyperthyroidism (Fig. T.13)
  - Secondary hyperthyroidism
• Goitre of thyroid deficiency:
  – Congenital hypothyroidism (cretinism)
  – Hypothyroidism (myxoedema)
  – Lymphadenoid goitre (Hashimoto's disease)
  – Drugs (e.g. resorcinol, phenylbutazone)
• Malignant enlargement:
  – Carcinoma
    ○ Sarcoma (rare)

These conditions can be regrouped for diagnostic purposes as follows:
• Thyroid enlargement with hyperthyroidism:
  – Primary hyperthyroidism (diffuse enlargement)
  – Secondary hyperthyroidism:
    ○ Localized enlargement: toxic adenoma (rare)
    ○ Generalized enlargement: nodular goitre in which one nodule may be so large as to suggest a solitary adenoma, occasionally parenchymatous or even malignant goitre
• Thyroid enlargement with signs of deficient secretion:
  – Congenital:
    ○ Congenital hypothyroidism
  – Acquired:
    ○ Hypothyroidism (myxoedema: mild deficiency may be exhibited by colloid or malignant goitre)
    ○ Hashimoto's disease
• Thyroid enlargement: uncomplicated
  – Localized enlargement: one large nodule in a small nodular goitre, cyst, adenoma, Riedel's disease (early stages), malignant disease (early stages)
  – Generalized enlargement: nodular colloid goitre, lymphadenoid goitre, Riedel's disease (late stages), malignant goitre (late stages).

**THYROID ENLARGEMENT WITH HYPERTHYROIDISM**

Hyperthyroidism is characterized by the presence of symptoms of hyperthyroidism from the onset of the disease. In secondary thyrotoxicosis, these symptoms develop after a simple goitre has been present for a variable period, often many years. The diagnostic points of each condition are listed in Table T.3.

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**Table T.3 Diagnostic points of hyperthyroidism**

<table>
<thead>
<tr>
<th></th>
<th>Primary hyperthyroidism</th>
<th>Secondary hyperthyroidism</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age of onset</td>
<td>Young</td>
<td>Middle-aged</td>
</tr>
<tr>
<td>Onset</td>
<td>Acute</td>
<td>Insidious</td>
</tr>
<tr>
<td>Thyroid swelling</td>
<td>Not present before onset. Generalized soft elastic and vascular swelling, enlargement not gross May harden if iodine has been given</td>
<td>Present before onset. Enlargement may be considerable; frequently nodular</td>
</tr>
<tr>
<td>Exophthalmos</td>
<td>Generally present, often gross</td>
<td>Rare and, if present, slight</td>
</tr>
<tr>
<td>Heart</td>
<td>Tachycardia, but fibrillation and heart failure not common except in late or severe cases</td>
<td>Tachycardia. Cardiovascular failure most prominent symptom. Atrial fibrillation fairly common</td>
</tr>
<tr>
<td>Tremor and general excitability</td>
<td>Marked</td>
<td>Slight</td>
</tr>
<tr>
<td>Loss of weight</td>
<td>Marked</td>
<td>Present, not so marked</td>
</tr>
<tr>
<td>Increased perspiration</td>
<td>Marked</td>
<td>Present, not so marked</td>
</tr>
<tr>
<td>Results of iodine medication</td>
<td>Often striking improvement</td>
<td>Improvement, but of a lesser degree</td>
</tr>
<tr>
<td>Thyroid function tests</td>
<td>Raised</td>
<td>Raised</td>
</tr>
<tr>
<td>Radioidine uptake</td>
<td>Raised</td>
<td>Raised</td>
</tr>
</tbody>
</table>
Various eye signs are described in connection with exophthalmos, of which the following are the best known:

- von Graefe’s sign: lagging behind of the upper lid as the patient looks downwards
- Dalrymple’s sign: retracted lids causing a wide palpebral opening
- Stellwag’s sign: diminished frequency of blinking
- Möbius’ sign: an inability to maintain convergence for close vision

Dalrymple’s sign is fairly constantly present but may be found in other conditions, while the other signs are neither constantly present nor confined to exophthalmic goitre. Indeed, lid retraction alone may be found without true exophthalmos.

CONGENITAL HYPOTHYROIDISM

Usually, the thyroid is atrophic in this condition, but a goitre is occasionally present, especially in a long-standing case. An untreated patient is easily recognizable but one is seldom seen nowadays. Slow development, either physical or mental, should rouse a suspicion of thyroid deficiency, remembering other possible causes of backward development such as rickets, renal rickets and achondroplasia. The diagnosis of congenital hypothyroidism will not be detailed further as it is barely relevant.

HYPOTHYROIDISM

As in congenital hypothyroidism, the thyroid is only occasionally enlarged, and here again a detailed account will not be given. The characteristic symptoms of hypothyroidism include slowed mentality, coarse features, dry skin, brittle nails and sparse coarse hair, and a gain in weight, often gross.

Certain drugs (e.g. resorcinol as an external application, and phenylbutazone by mouth) may occasionally be associated with thyroid enlargement, with signs of hypothyroidism reversible on stopping drug administration. Occasionally, a moderately enlarged gland may increase in size during treatment with an anti-thyroid agent (e.g. neomercazole).

LYMPHADENOID GOITRE (HASHIMOTO’S DISEASE)

In this disorder, the thyroid gland becomes infiltrated with lymphoid tissue as a result of an autoimmune reaction. Interestingly, it was the first autoimmune disease to be described. It is a disease occurring in women in middle life, and usually produces a uniform, firm enlargement of the thyroid with evidence of hypothyroidism. Laboratory tests for thyroid function show low levels. There is an increased serum cholesterol, a raised erythrocyte sedimentation rate, and autoimmun antibodies are present in the blood.

Occasionally, lymphadenomatous goitre will occur with normal thyroid function and, very exceptionally, with hyperthyroidism. In the past, diagnosis has often been made after operation as a result of histological section of the tissue removed, but with careful investigation this should not be necessary in a typical case.

Carcinoma of the thyroid may cause confusion, but this condition is practically never associated with hypothyroidism in an untreated case.

RIEDEL’S DISEASE

This is an interesting and rare condition in which an intense sclerosing fibrosis starts in one area of the gland and spreads first to the whole gland, and then to surrounding structures. Its aetiology is unknown, but it may represent an end stage of Hashimoto’s disease. The progress is slow as a rule, but gradually the trachea, the oesophagus and the great vessels all suffer from constriction, while the recurrent laryngeal nerves are affected early. The differential diagnosis from malignant disease is very difficult, but the condition should be suspected when an intensely hard goitre with pressure symptoms out of all proportion to its size is found in a young adult. Diagnosis is confirmed by histological examination of biopsy material.

UNCOMPPLICATED THYROID ENLARGEMENT

A true adenoma is an uncommon condition, but a particularly large nodule in an otherwise small nodular goitre forming an asymmetrical swelling in the thyroid tissue is common (Fig. T.14). The lesion may be either cystic or solid, but palpation is not always reliable in determining this. Nodular goitre may give rise to a uniform enlargement, as also may colloid goitre. This last condition may present a smooth surface, as is usually the case in parenchymatous enlargement. A simple cyst is quite common but it may suddenly enlarge from haemorrhage into it.

Figure T.14 A large colloid mass in the right lobe of the thyroid gland has been outlined with a skin pencil.
Malignant disease starts in one area and spreads to involve the whole gland, finally breaking through the capsule to invade the surrounding structures. Movement on deglutition may be lost, the recurrent laryngeal nerve is involved early, and the growth tends to surround the carotid sheath rather than push it back, as is the case with large simple goitres, so that pulsation of this vessel may be impalpable in the middle of the neck. The sympathetic chain is often involved late in the disease, with a resultant Horner's syndrome. The swelling is usually hard, as in Riedel's disease, but it tends to be much greater in size and more rapid in growth. Pressure symptoms are early, and pain is often a marked feature, particularly on swallowing. Bone and lung metastases are not uncommon.

One type of thyroid carcinoma – namely the papillary carcinoma – deserves special mention. This lesion typically occurs in the fourth and fifth decades of life, when it metastasizes to the lymph nodes. The secondary deposits may be much larger than the primary, which cannot be detected, so that these cases often present with soft lumps in the side of the neck that used (erroneously) to be called 'lateral aberrant thyroids'. Thyroid tissue so situated is always a secondary deposit from a small primary in the thyroid.

LABORATORY INVESTIGATIONS
These include the following:

- **Serum free thyroxine (T4) and free tri-iodothyronine (T3):** Measurement of the biologically active unbound fraction is more accurate than measurement of total T3 and T4; elevation suggests hyperthyroidism.
- **Thyroid-stimulating hormone (TSH) level:** Raised in myxoedema, but suppressed in hyperthyroidism, where the gland secretes T4 autonomously.
- **Thyroid scintogram:** Radiiodine studies of the thyroid gland provide very useful information. A small tracer dose of gamma-ray-emitting iodine-131 is injected intravenously and the gland scanned with a gamma-ray detector to map areas of high uptake reflecting high activity. A nodule in the thyroid gland that is hyperactive can be pinpointed by this method, a so-called 'hot nodule'. Similarly, a nodule that is not producing T4 will not take up the radiiodine, for example a cyst or tumour ('cold nodule').
- **Thyroid antibodies:** Against thyroglobulin or the 'microsomal' antigen (now identified as thyroid peroxidase), indicate an autoimmune pathology (Hashimoto's thyroiditis); other autoantibodies are often present.
- **Thyroid ultrasound:** Provides valuable information on whether a mass is solid or cystic.
- **Fine-needle aspiration and biopsy:** Allows material to be obtained for cytological and histological examination. It is now the principal investigation for all solitary nodules, often under ultrasound guidance.
- **Serum cholesterol:** Usually raised in myxoedema and may be normal or a little low in hyperthyroidism.
- **Electrocardiogram:** In hyperthyroidism cardiac involvement will show low electrical activity with small complexes. Atrial fibrillation complicating hyperthyroidism will be confirmed.

THYROID, PAIN IN
Harold Ellis

A painful thyroid swelling is not a common clinical situation. The two most likely causes to be encountered are either haemorrhage into a pre-existing thyroid cyst or inflammation of a thyroglossal cyst. The following conditions may give rise to this pain symptom:

- **Inflammatory:**
  - Acute (suppurative) thyroiditis
  - Subacute thyroiditis (De Quervain’s disease)
  - Inflammation of a thyroglossal cyst or fistula
- **Haemorrhage into a cyst of the thyroid**
- **Hashimoto’s disease (rarely)**
- **Carcinoma of the thyroid in its late stages**

ACUTE (SUPPURATIVE) THYROIDITIS
This is a rare condition that is nearly always bacterial in origin. Fungal and parasitic causes can be regarded as medical oddities. The usual organisms producing this condition are *Staphylococcus aureus*, haemolytic *Streptococcus*, pneumococcus, and occasionally *Salmonella* and *Escherichia coli*. In two-thirds of cases, there is pre-existing thyroid disease. The sexes are equally affected.

The source of the bacterial invasion is either extension from an adjacent infection or bacteraemia secondary to a distant focus. Commencing as an acute inflammation, the condition usually progresses to suppuration. The clinical features are a sudden onset with severe pain in the neck, which may be referred to the ear, the lower jaw or the occiput and which is aggravated by swallowing and movement of the neck. There is associated malaise and fever.

Examination reveals a febrile patient (the temperature in the range of 38–40 °C), and tachycardia. Swelling, tenderness and redness in the region of the thyroid generally appear later and, more characteristically, only one lobe of the thyroid is involved. Regional lymphadenopathy is variable. The neck is held...
flexed, and neck movement is painful. Fluctuation is not usually elicited because of induration of the surrounding tissues. There is leucocytosis and, if untreated, the mass progresses to the formation of an obvious abscess.

SUBACUTE (NON-SUPPURATIVE) THYROIDITIS
This condition, also known as De Quervain’s thyroiditis, is viral in origin. Any age may be affected, ranging from 3 to 76 years, although it occurs most commonly in the fifth decade of life. Females are far more often affected than males. Usually, the condition involves a previously normal gland.

The illness is preceded frequently by an upper respiratory infection, and the thyroid symptoms are often antecedent by muscular aches and malaise with fever (in the region of 39 °C) and weight loss. Pain then develops in the thyroid gland and possibly radiates to the ears. This is aggravated by movement of the neck and swallowing. Usually, both lobes are enlarged, although in one-third of cases one lobe is involved first, and the inflammation then spreads to the opposite side. Examination of the neck reveals a tender, firm or hard enlargement of the thyroid gland. Quite often there are accompanying symptoms and signs of hyperthyroidism.

Laboratory tests reveal a raised white cell count and erythrocyte sedimentation rate. Usually, the level of thyroxine is raised, the elevation persisting for 1–3 months.

The condition runs a variable course of weeks or months and, even if untreated, it usually subsides without sequelae. Rarely, it is followed by clinical hypothyroidism. As its name implies, it does not proceed to frank suppuration.

INFLAMMATION OF A THYROGLOSSAL CYST OR FISTULA
The typical thyroglossal cyst lies in the midline of the neck, usually at the cricothyroid space, less commonly at a higher or lower level, although it may deviate somewhat to one or other side of the midline. It usually presents in children or young adults, and it characteristically moves upwards on protrusion of the tongue, as well as on swallowing (see NECK, SWELLING OF, p. 454). Infection of the cyst is not uncommon, when it presents as an obvious inflammatory mass above the anatomical region of the thyroid gland. A thyroglossal fistula is occasionally congenital, but it may follow infection or inadequate removal of a thyroglossal cyst. The fistula discharges mucus and is frequently the site of recurrent attacks of inflammation.

Figure T.15 An advanced carcinoma of the thyroid. The sternocleidomastoid muscle is involved, and the cervical lymph nodes are enlarged and hard.

THYROID CYST
Haemorrhage into a pre-existing thyroid cyst produces a sudden, painful enlargement of a lump in the thyroid gland, which may or may not have already been noted by the patient. The danger is that it may also produce a sudden and dangerous compression of the trachea, with respiratory obstruction. The symptoms may require urgent surgical treatment but, if less severe than this, the swelling gradually subsides over the succeeding few days.

The cystic nature of the swelling can be confirmed by ultrasound examination of the mass.

HASHIMOTO’S DISEASE
This condition (see p. 680) is usually painless, but from time to time the thyroid enlargement may be painful and tender.

CARCINOMA OF THE THYROID
Poorly differentiated (anaplastic) carcinomas of the thyroid usually occur in elderly patients. In their advanced stages, the lesions produce a tender and painful infiltrating mass in the neck, usually with local lymphadenopathy (Fig. T.15). Clinical diagnosis is not usually in doubt, but it can be confirmed by needle biopsy.

TICS
Mark Kiniron

(See also TETANY, p. 671; VERTIGO, p. 736.)
‘Tics’ and ‘habit spasms’ are the terms that cover a variety of twitching or jerking movements that occur irregularly and tend particularly to involve the muscles around the eyes, the face and the shoulders. These are voluntary movements, and patients usually obtain
relief from tension by their repetitive performance. In many instances, the condition is exaggerated by anxiety or neurosis. Often, with the passage of time, the movements become so habitual as to be almost involuntary.

In most instances, the movement is the same and repeated in an identical fashion. Multiple tics may occur in the syndrome of Gilles de la Tourette. In this syndrome, the multiple tics may be accompanied by involuntary utterances and grunts. Although tics in isolation are thought to be functional in nature, there is some evidence to suggest that this particular syndrome has an organic neural basis. Multiple tics need to be differentiated from involuntary movement disorders such as chorea.

**TINNITUS**

Michael Gleeson

Tinnitus is a symptom that can be defined as any sound perceived by the patient when no external source of the sound exists. In the past, tinnitus was subclassified as either objective or subjective, depending on whether it was audible to the physician or not. However, objective tinnitus is extremely rare – albeit fascinating for the physician and alarming for the patient. A more appropriate and clinically useful classification is provided in Box T.6.

From a clinical perspective, there are two types of tinnitus that demand careful investigation:

- **Pulsatile tinnitus**, which may indicate the presence of a vascular abnormality or tumour
- **Unilateral tinnitus**, in which a space-occupying lesion within the cerebellopontine angle or petrous apex must be excluded

In all other respects, the quality of the tinnitus – no matter how disturbing for the patient – has little diagnostic relevance.

Tinnitus is a very common symptom, which most people experience from time to time. Most normal people perceive some tinnitus if placed in a completely silent environment. The majority of people manage to suppress the tinnitus from conscious awareness; this is compensated tinnitus. A few are unable to suppress their tinnitus and, regardless of its intensity, find that it interferes with every second and activity of their life; this is uncompensated tinnitus.

All patients with tinnitus should undergo a careful general and otological/clinical examination in order to exclude any pre-existing conditions that may suggest the underlying cause. Most important, a thorough inspection of the ears, nervous system and neck, together with pure tone audiometry, should be undertaken. Only when all underlying conditions have been excluded can the patient be considered to have idiopathic tinnitus.

Unilateral tinnitus always demands further investigation. It is a common and early presenting feature of VIIIth nerve tumours, and is not necessarily accompanied by hearing loss or vertigo. Patients with unilateral tinnitus should always have a magnetic resonance scan of their cerebellopontine angle.

Pulsatile tinnitus usually suggests a vascular abnormality although it is also a feature of benign intracranial hypertension. Light pressure on the neck over the internal jugular vein may abolish tinnitus in those patients with venous turbulence. An audible murmur on auscultation over the carotid artery or mastoid, or a palpable thrill, may suggest an arteriovenous malformation or carotid stenosis that can be confirmed by arteriography. Pulsatile tinnitus is the most common presenting symptom of patients with glomus tumours of the middle ear or skull base (Figs T.16 and T.17). These paraganglia are not always visible or palpable on clinical examination, and again, a magnetic resonance scan is necessary for diagnosis and carotid angiography for management planning.

If there is an obvious cause in the external or middle ear (foreign bodies, wax impaction, otitis externa or otitis media), treatment of this cause may eradicate or reduce the intensity of the tinnitus. In those patients in whom the tinnitus is an accompanying feature of a significant hearing loss, it may be improved by fitting an appropriate hearing aid.

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**Box T.6 Causes of tinnitus**

<table>
<thead>
<tr>
<th>Otological causes</th>
<th>Neurological causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>• External ear</td>
<td>• Ménière’s disease</td>
</tr>
<tr>
<td>– Otitis externa</td>
<td>– Ototoxicity</td>
</tr>
<tr>
<td>– Foreign bodies</td>
<td>– Skull base fracture</td>
</tr>
<tr>
<td>– Wax</td>
<td></td>
</tr>
<tr>
<td>• Middle ear</td>
<td>• Myoclonus of the middle ear muscles</td>
</tr>
<tr>
<td>– Otosclerosis</td>
<td>– VIIIth nerve tumours</td>
</tr>
<tr>
<td>– Chronic suppurative otitis media</td>
<td>– Temporal lobe epilepsy</td>
</tr>
<tr>
<td>– Acute otitis media</td>
<td>– Palatal myoclonus</td>
</tr>
<tr>
<td>– Tympanosclerosis</td>
<td>– Multiple sclerosis</td>
</tr>
<tr>
<td>– Middle ear tumours</td>
<td>– Glomus jugulare</td>
</tr>
<tr>
<td>• Inner ear</td>
<td>– Glomus tympanicum</td>
</tr>
<tr>
<td>– Presbyacusis</td>
<td>– Arteriovenous malformations</td>
</tr>
<tr>
<td>– Noise-induced hearing loss</td>
<td>– Carotid artery stenosis</td>
</tr>
<tr>
<td>– Idiopathic sensorineural hearing loss</td>
<td>– Jugular bulb turbulence</td>
</tr>
<tr>
<td>– Labyrinthitis</td>
<td>(venous bands)</td>
</tr>
</tbody>
</table>

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Rarer and treatable causes of tinnitus include palatal myoclonus and spasms of the middle ear muscles. Clenching the teeth often intensifies or changes the quality of the tinnitus in this group of patients. Botulinum toxin injection or section of the tensor tympani or stapedius tendons can prove effective.

In the majority of patients, the precise cause of their tinnitus is not immediately obvious. The cause is then usually attributed to minor degrees of age-related hearing loss. It must be remembered that tinnitus is an extremely disturbing symptom for the patient. Many become convinced that they have a brain tumour, or even something worse. An explanation of their symptoms, together with sensitive reassurance, allows many to live happily with their tinnitus.

Tinnitus maskers have limited benefit for a minority of sufferers, and trials of various medications have met with only minor success.

**TONGUE, DISCOLORATION OF**

Simon Anderson

The mucous membrane of the tongue is covered mainly by filiform papillae, which cover the major portion of the tongue, and vary in length from 1 to 3 mm. Fungiform papillae are found at the apex and along the lateral aspect of the tongue. They are barely visible by eye, but on occasion, even in the healthy individual, they may be red, large, smooth and round.

Today, discoloration of the tongue has much less of a diagnostic role in gastrointestinal disease than it did in the past. Changes that were thought to take place with constipation, appendicitis and other gastrointestinal diseases are insignificant and unreliable. On the other hand, there are more important changes that are associated with infectious diseases, deficiency states and metabolic disorders.

The tongue surface may be dry, brown and slightly furred in patients who are mouth-breathers or who are dehydrated. A brown, dry tongue is common in smokers. Furring of the tongue is due to heaping up of the squamous epithelium on the filiform papillae, from inadequate cleansing of the tongue. When this is more pronounced, a black hairy tongue is seen (see Fig. M.22) with very marked overgrowth of the filiform papillae in patients on antibiotic therapy, smokers and those with poor oral hygiene.

Hyperpigmentation of the tongue (a black tongue) is commonly seen in dark-skinned individuals. This can also be seen with certain medication such as tetracyclines, bismuth, antidepressants and interferon. Pigmentation of the tongue may be found in Addison’s disease and acanthosis nigricans, in which the tongue undergoes hypertrophy of the filiform papillae to produce a shaggy, papillomatous dorsum.

The term ‘geographic tongue’ (benign migratory glossitis) is used when an area of filiform papillae is lost from the dorsum of the tongue. Smooth, pink mucosa is seen that contrasts sharply against that mucosa which is covered with normal papillae. A feature of the condition is that the appearance of the tongue will change from day to day, creating a ‘wandering rash’ across the surface of the tongue. This condition is benign and can be regarded as a variant of normal.

Hypertrophy of the fungiform papillae in scarlet fever (streptococcal pharyngitis) produces the classic ‘strawberry’ or ‘raspberry’ tongue.

**Figure T.16** Typical otoscopic appearance of glomus tympanicum. The vascular mass is readily visible behind the eardrum.

**Figure T.17** Internal carotid angiogram showing the intense vascular blush produced by a large jugular paraganglioma.
A fissured tongue is a benign condition characterized by deep grooves in the middle of the tongue. It is seen in Down’s syndrome and the Melkersson–Rosenthal syndrome. The cause is unknown and is not associated with other underlying disease. Orofacial granulomatosis is an idiopathic granulomatous inflammation of the lips and mouth and also results in a fissured tongue. Leucoplakia describes white patches on the surface or lateral aspect of the tongue. These, on histology, show hyperkeratosis, acanthosis and dyskeratosis. The white areas may be seen as a patch or a raised plaque, and are usually painless. Leucoplakia is regarded as a precancerous lesion, as is the asymptomatic red, velvety lesion that may sometimes be seen on the ventrolateral aspect of the tongue. ‘Hairy’ leucoplakia is unique to HIV infection and is found particularly along the side of the tongue, although it may occur anywhere in the mouth. The lesions are slightly raised and poorly demarcated, and show a corrugated white ‘hairy’ surface that does not rub off; they are asymptomatic. The histology is distinctive, showing keratin projections, parakeratosis, acanthosis and a characteristic ballooning change in the prickle-cell layer.

Mucosal infection with the yeast Candida albicans produces the clinical picture of moniliasis (candidosis or thrush). Creamy white, curd-like patches occur on the tongue and other areas of the buccal mucosa. When scraped, they reveal a raw bleeding area. The lesions can be quite painful and are found particularly in sick infants, debilitated patients or those receiving broad-spectrum antibiotics or high-doses of corticosteroids. Candidiasis is a common manifestation of immunodeficiency, particularly HIV infection. In infants, thrush is distinguished from milk curds by the difficulty with which the former lesions are removed, leaving an underlying patch of infected mucosa. Oral candidiasis can also cause red lesions without white plaques – erythematous ‘atrophic’ candidiasis. It can also be a sign of infection elsewhere – in the oesophagus, angular cheilitis or sytemic infection.

The tongue will appear blue in patients who are centrally cyanosed, and pale in anaemic patients. Many anaemic patients, however, have atrophic glossitis – a red, painful, featureless tongue. This is the consequence of atrophy of the papillae, which may occur in pernicious anaemia, severe iron-deficiency anaemia and/or deficiency states involving other B vitamins. The red, painful tongue may be associated with other evidence of mucosal atrophy in the mouth and fissuring at the corner of the lips known as ‘angular stomatitis’ or cheilitis. The mucosal lesions of folic acid deficiency are often more marked than those encountered in vitamin B12 deficiency. Other terms that have been used to describe the colour changes taking place in vitamin B deficiencies are the ‘beefy red’ tongue of pellagra and the ‘magenta’ tongue of riboflavin deficiency. The Plummer–Vinson (or Patterson–Kelly) syndrome describes the iron-deficiency state in which the tongue is painful, reddened and smooth. There is cheilitis, postcroroid webs and koilonychia. Acrodermatitis enteropathica is a rare autosomal recessive disorder related to zinc deficiency. Intraoral features include a white coating to the tongue and buccal mucosa, with marked halitosis. There is chronic diarrhoea, hair loss, severe dermatitis and failure to thrive. The condition may also occur in patients on maintained hyperalimentation who become deficient in zinc.

In hereditary haemorrhagic telangiectasia (Osler–Weber–Rendu syndrome), telangiectasia occur throughout the gastrointestinal tract. They most commonly occur in areas of the oral cavity, including the lips, gingiva, buccal mucosa and tongue. The lesions are dilated capillary vessels and small arterioles. The pigmeny changes involving Peutz–Jegher’s syndrome and pseudoxanthoma elasticum do not normally involve the tongue. Granulomatous involvement of the tongue and buccal mucosa in Crohn’s disease produces raised, smooth, red nodules and hyperplastic ridges on the tongue. These may appear erythematous. A number of dermatological diseases may affect the mouth, including erythema multiforme, pemphigus and pemphigoid, but involvement of the tongue in these conditions is rare. Lichen planus may occur only in the mouth or as part of the more common cutaneous and genital involvement. The features vary from lace-like white patches (Wickham striae) on the buccal mucosa to painful gingival erosions.

## TONGUE, PAIN IN

Harold Ellis

Pain in the tongue may be attributable to an obvious lesion, usually with breach of the surface, such as a carcinoma. These conditions are discussed elsewhere (see TONGUE, ULCERATION OF, p. 689). On the other hand, pain in the tongue or soreness of the tongue may be an insistent complaint when there is no superficial abnormality. The conditions that must be considered include the following:

- When the pain complained of is not on the dorsum, tip or sides of the tongue but underneath or deeper:
  - Injury to the frenulum linguæ
  - Ranula
TONGUE, PAIN IN

- Calculus in the duct of a submandibular salivary gland
- Foreign body in the tongue
- Myositis
- Trichinosis

- When the pain complained of appears to be upon the surface of the tongue, even if it also affects the tongue as a whole:
  - Bitten tongue
  - After an anaesthetic (mouth-gag)
  - Injury by tooth or dental plate
  - Antibiotic glossitis, associated with lichen planus, Behçet’s disease or pemphigus vulgaris
  - Congenital fissured tongue
  - Geographic tongue
  - Median rhomboid glossitis
  - Moeller’s glossitis
  - Glossitis of deficiency disease
  - Smoking
  - The effects of over-hot beverages or foodstuffs
  - The effects of pungent condiments such as cayenne pepper
  - Minor viral diseases
  - Carcinoma

The differential diagnosis depends upon the following considerations.

PAIN UNDERNEATH THE TONGUE, OR DEEPER

Injury to the frenulum linguae may cause visible abrasion or a definite ulcer. The most injured spot is tender as well as painful, the diagnosis depending on careful attention to the appearance and to the site of greatest tenderness. The cause may be injury by a fish bone or other sharp or puncturing objects. In violent coughing bouts, as in whooping cough, the protruded tongue may be forced against the lower incisor teeth with such violence that the frenulum becomes abraded, inflamed or ulcerated.

A ranula is not painful unless it becomes inflamed. It is an asymmetrical, red, smooth cystic swelling in the floor of the mouth under the tongue on one or other side of the frenum. It may result from obstruction of the duct of one of the sublingual salivary glands, but more often it is a retention cyst arising in one of the many mucous glands in the floor of the mouth.

Calcium in the duct of a submandibular salivary gland is not necessarily painful. It may produce discomfort or more or less severe pain, recurrent or constant according to the degree of inflammation. The stone may be very small and difficult to detect either with a probe or by X-rays. However, its existence may be suspected by the situation of discomfort, or by the corresponding salivary gland swelling when the patient begins to eat, when the stone interferes with the free passage of increased saliva flow. The calculus can frequently be palpated bimanually in the floor of the mouth, and it is occasionally seen to protrude through the duct orifice. An X-ray will confirm the diagnosis (see Fig. S.3).

A foreign body in the tongue is uncommon, although a fish bone may become impacted in it. More often, the foreign body injures the tongue, itself escaping but leaving the pain behind. The diagnosis depends on the accuracy of the story obtained, or the discovery of the foreign body by palpation or by radiography.

Myositis of the tongue is seldom (if ever) a localized condition. It may, however, be a prominent feature in polymyositis or in trichinosis, in which the embryo trichinellae have a special predilection for the muscles at the base of the tongue, which become stiff, painful and tender. The diagnosis of trichinosis is difficult, especially as it will hardly be thought of unless there is an epidemic at the time. The blood exhibits eosinophilia, but the only way of clinching the diagnosis is by demonstrating the trichinellae embryos microscopically in portions of the muscles excised.

PAIN ON THE SURFACE OF THE TONGUE

A bitten tongue will usually present an obvious lesion, but the pain may persist after a tongue bite, even when no obvious bruising or breach of the surface can be detected. The patient may be unaware of having accidentally inflicted the bite, notably if the accident occurred during sleep or during an epileptic seizure. Indeed, the occurrence of a local painful area in the tongue suggesting the effect of tongue bite may be the first indication that the patient has epilepsy. In tetanus, traumatic glossitis is common and may cause airway obstruction. After general anaesthetics, patients may complain of soreness of the tongue resulting from the use of tongue forceps or of a mouth-gag.

Injury by a tooth or dental plate may cause a local painful place on one side of the tongue, often fairly far back, the pain being increased by movements of the tongue in speaking, eating or swallowing. Fear of cancer is usual until the cause is found in the jagged edge of the adjacent tooth, or of the dental plate at the corresponding site. The condition needs to be watched carefully to be certain that the lesion disappears after the offending irritant is smoothed down or removed, and to allay any anxiety that the jagged tooth or plate may have initiated an epithelioma. Tuberculosis of the tongue, presenting as a painful deep persistent ulcer, is now rarely seen.
Antibiotic glossitis is a common cause of diffuse soreness of the tongue from the taking of antibiotics by mouth. The pain is sometimes due to infection with *Monilia albicans*, which can be grown from the surface. Its preponderance is favoured by the wide-spectrum antibiotics. In other cases of antibiotic glossitis, no such cause can be found, and the change is attributed to vitamin deficiencies arising from the suppression of normal gut flora. The tongue is clean, red and very sensitive to heat (Fig. T.18). Glossitis occurs in deficiencies of vitamin B12 and folic acid, in pellagra, malabsorption syndrome and Plummer-Vinson syndrome, but it seldom causes acute pain in these conditions.

*Lichen planus* affecting the tongue may be confused with monilial glossitis because both conditions produce small whitish patches on the surface. The lichen tends to produce lines or a mesh of pearly dots, and to favour the cheeks near the occlusal line of the molars.

The tongue may also become inflamed and painful in Behçet’s disease, erythema multiforme or pemphigus vulgaris.

*Congenital fissured tongue* (Fig. T.19) or ‘scrotal tongue’ is thick, deeply fissured and usually symptomless. If food particles lodge in the fissures, infection may arise and cause pain.

*Geographic tongue* (Fig. T.20) shows red denuded patches of irregular outline, which often change their position. It causes anxiety rather than pain.

*Median rhomboid glossitis* (Fig. T.21) is a rare congenital abnormality caused by persistence of the tuberculum impar between the two halves of the tongue. It occupies the middle third of the dorsum and is smooth, shiny and red. It carries
TONGUE, SWELLING OF

Moeller’s glossitis, often confused with the glossitis of pernicious anaemia, presents atrophic sharply defined red patches on the dorsum and sides: the atrophy in pernicious anaemia is evenly spread, and the mucosa is pale and dry. Spiced food causes pain. The condition may be met in allergic states, in nutritional deficiencies and with certain drug eruptions (e.g. reserpine).

Glossitis of deficiency disease occurs with avitaminosis, particularly of the B group, as in pellagra, but also with iron-deficiency and pernicious anaemia.

Smoking and the effects of tea or other hot liquid or food may cause acute pain in the tongue lasting for days after the cause has ceased to act. Pungent condiments such as capsicum, cayenne pepper, ginger and the like may similarly be responsible.

Foot-and-mouth disease may rarely be contracted by humans from infected farm animals, or from consumed milk or milk products from infected herds. Vesicles appear in the mouth and on the tongue. In the so-called ‘hand-foot-and-mouth disease’, which is probably due to Coxsackie A viruses, children are affected. Vesicular stomatitis contracted from horses, cattle and pigs occurs in North and South America.

Carcinoma of the tongue (Fig. T.22) starts as a nodule, fissure or ulcer, usually on the lateral border of the organ. At first painless, it becomes painful as it invades and becomes grossly septic. The pain often radiates to the ear, being referred from the lingual branch of the trigeminal nerve supplying the tongue along its auriculotemporal branch. Ulceration is accompanied by bleeding; hence the typical picture of late disease is an old man spitting blood into his handkerchief with a plug of cotton-wool in his ear.

**TONGUE, SWELLING OF**

Harold Ellis

Swelling of the tongue is a condition the nature of which is generally obvious on inspection and palpation, if the history is taken into account at the same time. Causes of tongue swelling are listed in Box T.7, although many require little detailed discussion.

If the nature of the tongue enlargement is not obvious from the history and simple inspection and palpation – as will probably be the case when it is due to a bite, sting, injury, corrosive or irritant application, after the use of serum, mercury, aspirin or other drugs, or in variola, pemphigus or erythema multiforme – it may be so from the concomitant symptoms, as in the case of congenital hypothyroidism (cretinism), acromegaly, Down’s syndrome or myxoedema.

Simple macroGLOSSIA is rare; when it does occur, the history is that it dates from youth or childhood, and the patient may otherwise be perfectly normal, unless he

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**Figure T.22** Carcinoma of the tongue.

<table>
<thead>
<tr>
<th>Acute swelling</th>
<th>Chronic or persistent swelling</th>
</tr>
</thead>
<tbody>
<tr>
<td>• Bite or sting</td>
<td>• Macroglossia</td>
</tr>
<tr>
<td>• Injury (e.g. by a fish bone, or by biting during an epileptic fit)</td>
<td>• Congenital hypothyroidism (cretinism)</td>
</tr>
<tr>
<td>• Corrosives or acute irritant applications</td>
<td>• Myxedema</td>
</tr>
<tr>
<td>• Acute oedema, secondary to:</td>
<td>• Mongolism</td>
</tr>
<tr>
<td>– Inflammatory conditions within the mouth: stomatitis</td>
<td>• Acromegaly</td>
</tr>
<tr>
<td>– The effects of certain drugs (e.g. mercury, rarely aspirin)</td>
<td>• Primary amyloidosis</td>
</tr>
<tr>
<td>– Erythema bullousum or pemphigus</td>
<td>• Local or asymmetrical swelling</td>
</tr>
<tr>
<td>– Varioila</td>
<td>– Irritation of a dental plate or decayed tooth</td>
</tr>
<tr>
<td>– Serum injections and other conditions liable to cause giant urticaria</td>
<td>• Carcinoma</td>
</tr>
<tr>
<td>– Angioneurotic oedema (angio-oedema)</td>
<td>• Gumma</td>
</tr>
<tr>
<td>• Haemorrhage into the substance of the tongue (e.g. in scurvy, leukaemia and other haemorrhagic states)</td>
<td>• Leucoplaikia (chronic superficial glossitis)</td>
</tr>
<tr>
<td>• Suprahyoid cyst</td>
<td>• Tuberculous infiltration</td>
</tr>
<tr>
<td>• Haemangiomia or lymphangioma</td>
<td>• Actinomycosis</td>
</tr>
<tr>
<td>• Sarcoma</td>
<td>• Ranula</td>
</tr>
<tr>
<td>• Lipoma</td>
<td>• Calculus in a sublingual salivary gland</td>
</tr>
<tr>
<td></td>
<td>• Mercury</td>
</tr>
</tbody>
</table>

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**Box T.7 Causes of swelling in the tongue**
or she also has some other congenital peculiarity, such as macrocheilia (blubber-lips).

The chronic local lesions associated with swelling are in many cases accompanied by superficial ulceration, and the difficulties that may arise in distinguishing simple, syphilitic and carcinomatous ulcers are discussed under TONGUE, ULCERATION OF see below. Tuberculous and actinomycotic glossitis are both rare and may be mistaken for malignant or syphilitic disease. Tuberculous lesions are usually painful, and this cause should always be thought of when considering the possible causes of a painful swollen tongue, particularly as the manifestations of tuberculosis of the tongue may assume unusual and bizarre forms. Ranula and sublingual salivary gland calculus or cyst both cause swellings that are beneath the front part of the tongue rather than in its substance, generally bulging up one side of the floor of the mouth near the frenulum linguae. A ranula is a distended mucous gland, and after growing to perhaps the size of a chestnut it often ceases to enlarge further; it does not fluctuate in its dimensions in relationship to meals, as a salivary gland swelling often does.

A suprathyroid cyst is situated in the root of the tongue posteriorly, when it arises from the remains of the embryological thyroglossal duct. It is seldom large; its nature is suggested by its situation.

An angioma of the tongue is rare (Fig. T.23). Sometimes, however, after remaining latent for years, it grows with rapidity and necessitates an operation. The diagnosis may be suggested by the colour of the tumour, but histological examination subsequent to removal may be required before one can be sure whether the tumour is a simple angioma, an angiosarcoma or a sarcoma.

A lipoma occurs infrequently in the tongue, and its lobulated form generally breaks the surface and presents like a cluster of soft white cherries.

Haemorrhage into the substance of the tongue, with swelling and inability to speak or eat, may result from certain blood disorders, such as acute leukaemia or primary or secondary thrombocytopenic purpura (see PURPURA, p. 554).

Acute oedema of the tongue may be due to severe stomatitis, angioneurotic oedema of the tongue or Ludwig’s angina. This is an acute inflammatory condition, often streptococcal in origin, affecting the floor of the mouth and tongue, and spreading rapidly through the deeper structures of the mouth, throat and neck, causing extreme swelling of the adjacent tissues.

Angioneurotic oedema of the tongue is rare, but it is important because it may, rarely, prove fatal. As a rule, there is a history of previous similar attacks in other parts of the body, and other members of the family may have had similar episodes. Tracheostomy may, although very rarely, be necessary as a life-saving measure, the diagnosis becoming clear only when the oedema of the tongue and adjacent parts subsides almost as rapidly as it came on, and the patient develops similar episodes (angio-oedema), probably in other parts, on subsequent occasions.

**TONGUE, ULCERATION OF**

Harold Ellis

To enable a good view to be obtained of the affected part, the patient should be seated in a good light, and the protruded tongue gently dried with a piece of gauze. The presence of an ulcer being ascertained, its nature may be considered under the following headings:

- Carcinomatous
- Syphilitic
- Dental
- Tuberculous
- Ulcer in connection with stomatitis

**CARCINOMATOUS ULCER**

Carcinomatous ulcer (see Figs T.22, T.25 and T.26) is much more common in men than in women. It is very unusual before the age of 30, and rarely starts before 45. The foul smell of the breath and the ill and wearied expression of the patient may awaken suspicion before the tongue is seen, as the sloughing ulcer is usually heavily infected, and the toxic absorption combined with pain and loss of sleep have a rapid and marked effect upon health.
The tongue in a normal individual can be protruded 3–4 cm beyond the teeth; if the protrusion is limited, or if the tongue is not protruded straight, it can generally be inferred – except in cases of paralysis – that there is some tumour binding it down (ankyloglossia). The position of the ulcer is to be studied, as is its relation to any sharp and carious tooth. A carcinoma is usually on the side of the tongue, but it may be anywhere on the upper, lateral or undersurface or on the floor of the mouth. It is hardly ever exactly in the midline, however.

As regards the ulcer itself, the typical appearance when fairly developed may be described as irregular, deep, foul, sloughy, with raised nodular everted edges and a surrounding area of induration. Other types associated with minimal ulceration of the mucous membrane are: the *scirrhous*, where there is an excessive fibroblastic reaction, and the affected part of the tongue is shrivelled up as in the similar atrophic scirrhous cancer of the breast; and the *nodular*, where the lesion is mostly buried within the substance of the tongue and, like an iceberg, broaches the surface over a deceptively small area. In addition, the papilliferous type and multiple ulcerations are not uncommon.

Lastly, there is the fissure carcinoma associated with *leucoplakia* (Fig. T.24) with thickened white patches of mucosa. This may have been predisposed to by smoking, syphilis, sepsis, spices, sore tooth or spirits, but often no cause can be found.

Except in early cases, some of the lymph nodes are enlarged and hard, and they may be fixed. The submandibular group is generally the first affected, but the disease sometimes misses these and invades the jugular and even the supraclavicular nodes. Examination, therefore, should not be concluded before the whole of the neck has been palpated. The diagnosis should have been made, however, before the disease has developed thus far. In its earliest stages, a carcinoma may be represented by a superficial ulcer no more than 1.5 cm in diameter, by a crack or a small lump, without any enlargement of the nodes. In all of these conditions, the ulcer is already hard and very resistant to any form of simple topical treatment. Any ulcer of the tongue occurring in a middle-aged man, and lasting for more than 2–3 weeks, should always awaken suspicion (Figs T.25 and T.26).

**Differential diagnosis**

*From syphilitic ulcer*

This may be a very real difficulty, owing to the fact that the two conditions may exist side by side, and that the syphilitic leucoplakia may be the actual precursor of a cancer. Positive serological reactions, therefore, are not
proof that a carcinoma is not present. If a well-formed
S. aureus is present, anti-syphilitic remedies soon make
a great change in its appearance. Although a biopsy
is necessary for a definite diagnosis, certain clinical
criteria are characteristic, and a putative diagnosis
of syphilis may be made when the ulcer is centrally
situated, aching, and with serpiginous in outline, and has
the steep-cut edges and wash-leather slough base
typical of syphilitic ulcers elsewhere.

From dental ulcer
The ulcer in this case is caused by a carious or
otherwise jagged tooth, and therefore is in a
corresponding position on the tongue. Furthermore,
the ulcer is soft to the touch, and heals rapidly when
the offending tooth is stopped or extracted. There is
seldom difficulty in differentiation, except when the
ulcer is of very long standing.

SYPHILITIC ULCER
This may be primary, secondary or tertiary.

Primary syphilis or chancre is certainly rare on the
tongue and, owing partly to its rarity and partly to the
fact that it is unexpected, it is frequently missed. It is
more common in men than in women, but it may
occur even in children. It starts as a small pimple
that ulcerates and becomes indurated, although the
induration is not so marked as when it is situated on
the glans penis. The appearance of a secondary rash
with general enlargement of the lymph nodes would
indicate the true diagnosis. Further proof is supplied
by positive serological tests and the detection of
spirochaetes in serum from the sore. Furthermore,
the sore heals rapidly under the influence of
treatment.

Secondary syphilis manifests itself by the formation of
mucous patches and superficial ulcers. The latter are
always almost multiple and situated along the edges
and tip of the tongue, and with them are also found
similar sores on the mucous membrane of the cheek,
lips, palate and tonsil, and at the edges of the mouth.
The ulcers are small, round, painful, with sharply cut
dges and a greyish floor. Other secondary symptoms
will be present to make the diagnosis clear.

Tertiary syphilis or gummatous ulcerations are now
extremely rare and are divided into superficial and
deep. Superficial gummases begin as small, round-
celled infiltrations in the mucous and submucous
tissue. The ulcers are usually shallow, often irregular,
and associated with chronic glossitis, fissures and
leukoplakia. Although rare today, they are extremely
important as they may be followed by carcinomatous
change. The ulcers themselves are not at first indurated
but, if they are surrounded by interstitial fibrosis,
many appear hard; a histological examination is
essential if there is the least doubt. A deep gumma
starts as a hard swelling in the substance of the
tongue. It is usually situated in the midline, and in the
posterior half. Later it softens, breaks down and shows
itself as a deep cavity with irregular soft, steep-cut
walls and a wash-leather-like slough at its base. It is
not painful and does not increase progressively in size.
The important thing is to distinguish it from carcinoma
and tuberculosis disease. Unlike carcinoma, it does not
infiltrate widely or fix the tongue, its history is short,
and it causes no pain. Furthermore, it yields rapidly to
anti-syphilitic treatment.

DENTAL ULCER
Dental ulcer is due to repeated small injuries from the
sharp edge of a decayed tooth or damaged denture
and is situated opposite the tooth, generally on the
side of the tongue. The ulcer is small, superficial
and not indurated, unless it is of long standing. It is
therefore not easily mistaken for any other kind of ulcer
but, if doubt arises, this can be allayed by the healing
of the ulcer on appropriate dental treatment. Failure to
heal within a fortnight suggests that it is a carcinoma.
There is a form of dental ulcer that is found on the
fraenum of the tongue in children suffering from
whooping cough. During the violent expiratory
spasms peculiar to the illness, the undersurface of
the tongue may suffer from rubbing over the lower
incisor teeth.

TUBERCULOUS ULCER
This is rare in the Western world, but it occurs at
that period of life during which tuberculous disease
of the lung is common, between the ages of 15 and
35 years. It is due to infection with tubercle bacilli
brought up into the mouth. The ulcer itself is usually
on the tip of the tongue or the side in its anterior
half, and is generally painful, although sometimes
it is entirely painless. The outline is irregular. The edges
are usually thin and undermined, and the base is
covered by pale granulations, or excavated
clearly down to the underlying muscle fibres. Less
commonly, the edges are raised, although never
evolved or hard, and the base is nodular, sloughy or
caseous. It has often been mistaken for a carcinoma
or gumma. The fact that it is not hard, that it is
usually painful, and that pulmonary tuberculosis is
present should point to the true diagnosis. Negative
serological tests exclude a syphilitic gumma,
although histological proof may be necessary by
biopsy and cultures carried out for bacteriological
confirmation.
A further example of tuberculous ulceration is the so-called ‘truncated tongue’. In this type, there is an oedematous infiltration of the parenchyma of the tongue, causing it to become swollen and almost ‘woody’. There is also a shallow ulceration of the tip, giving an appearance as if part of the tongue has been amputated. In fact, the clinical manifestations of tuberculosis of the tongue are so protean that this disease should always be suspected in unusual lesions, especially if associated with pain.

ULCERS IN CONNECTION WITH STOMATITIS (ULCERATIVE STOMATITIS)

Septic infection of the mouth due to a variety of causes, such as irritation from decayed teeth, alkalis, acids or mercury, may be accompanied by the formation of small vesicles, which, on bursting, give rise to superficial ulcers (Fig. T.27). They are not limited to the tongue, but also appear on the mucous membrane of the cheeks and gums. Aphthous stomatitis commonly occurs in conjunction with the febrile diseases of childhood. It is characterized by the formation of whitish spots on the buccal mucous membrane, and small superficial ulcers may be formed by shedding of the epithelium. The ulcers of the tongue here occur during the course of a general inflammation of the mouth. One type that may be resistant to treatment is produced by Vincent's angina organisms; bacteriological tests provide the diagnosis, but it may be suggested by the extreme foetor of the breath.

When ulceration of the tongue – and at the same time, very probably of the inside of the mouth in general – occurs in such conditions as chickenpox, pemphigus and other conditions that may affect the buccal mucosa as well as the skin, the diagnosis depends, not upon the appearances of the ulcers or the tongue, but upon the concomitant skin eruption.

TONSILS, ENLARGEMENT OF

Michael Gleeson

The tonsils are situated between the anterior and posterior faucial pillars of the oropharynx. They consist of lymphoid tissue and play an important role in the development of the immune system during childhood. There is a physiological increase in size of the tonsils between the ages of 4 and 6 years, and subsequently the tonsils involute. During childhood, tonsillar enlargement is usually accompanied by a synchronous enlargement of the adenoids. Extreme hyperplasia of the tonsils at this age (‘kissing tonsils’) can lead to respiratory obstruction and obstructive sleep apnoea. These patients improve dramatically with adenotonsillectomy.

In adult life, enlargement of the tonsils is most commonly caused by infectious diseases. Symmetrical enlargement may be caused by bacterial infection, glandular fever or toxoplasmosis, and it is then accompanied by severe malaise and lymphadenopathy.

A peritonsillar abscess (‘quinsy’) develops when infection spreads through the capsule of the tonsil (see Fig. T.7). Pus and oedema displace the tonsil medially and cause trismus, drooling, dysphagia and a change in quality of the voice (‘hot potato voice’). Dramatic relief of many of these symptoms is experienced when the pus is evacuated.

The head and neck is the most common region for the development of lymphoma, and many of these present as unilateral enlargement of the tonsil. This may, or may not, be accompanied by enlargement of the cervical lymph nodes.

In adults, deep lobe or parapharyngeal parotid tumours can cause displacement of the tonsil that can be mistaken for tonsillar enlargement. Other lesions in the parapharyngeal space, such as deep lobe parotid tumours, carotid and vagal paragangliomas or schwannomas, can cause the same tonsillar displacement and apparent enlargement of the tonsil.

TRACHEAL DEVIATION

Alex West

The trachea extends downwards from the lower border of the cricoid cartilage to the bifurcation into the two main bronchi at the level of the fifth thoracic vertebra. In normal individuals, the upper half of the trachea lies in the midline of the neck, while the lower intrathoracic portion inclines slightly to the right of the midline. Palpation of the trachea in the neck may provide
information about the position of the mediastinum, provided that the thoracic spine is straight and the thyroid gland not enlarged.

Displacement of the trachea from the mid-sternal line will usually signify disease of the pleura, lung or occasionally the mediastinum. In conjunction with conventional physical examination of the chest to assess chest wall movement, air entry, percussion note, vocal fremitus and breath sounds, the position of the trachea should be carefully palpated. Deviation of the trachea away from the abnormal side will suggest volume displacement, perhaps due to a large pleural effusion, pneumothorax under tension, bullous emphysema, or, rarely, a massive lung or mediastinal tumour (Fig. T.28). Deviation towards the abnormal side will suggest collapse of the underlying upper lobe or lung. In adults, deviation of the trachea towards the affected side in conjunction with flattening of the upper anterior chest raises the possibility of long-standing fibrotic contracture of the lobe or lung, possibly with associated pleural thickening, secondary to healed pulmonary tuberculosis.

The trachea is not displaced by consolidation without collapse of the underlying lung. It may remain centrally positioned when carcinoma has caused a combination of lung collapse with pleural effusion. A confirmatory chest radiograph should be taken if the physical signs suggest underlying lung or pleural disease.

**TREMOR**

David Werring & Mark Kinirons

Tremor may be defined as a regular, rhythmic oscillation of one part of the body from a fixed point, usually in one plane. It results from alternating or synchronous contractions of reciprocally innervated antagonistic muscles. Any body part, including the limbs, neck, tongue, chin or vocal cords, may be affected. There are three basic types of tremor: (i) a resting tremor is present with the relevant body part completely reposed; (ii) a postural tremor is elicited by extending a limb against gravity (this term is often used interchangeably with action tremor); and (iii) an intention tremor is elicited by moving a limb to and from a target. The differential diagnosis for each type is listed in Box T.8.

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**Box T.8 Types of tremor**

<table>
<thead>
<tr>
<th>Type of tremor</th>
<th>Causes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Resting tremor</td>
<td>- Idiopathic Parkinson’s disease</td>
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<tr>
<td></td>
<td>- Other parkinsonian syndromes (e.g. MSA-P)</td>
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<tr>
<td></td>
<td>- Essential tremor (resting tremor usually much less prominent than postural/intention tremor)</td>
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<tr>
<td></td>
<td>- Wilson’s disease</td>
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<tr>
<td>Postural tremor</td>
<td>- Physiological</td>
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<td>- Exaggerated physiological tremor</td>
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<td></td>
<td>- Stress/anxiety</td>
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<td></td>
<td>- Thyrotoxicosis, phaeochromocytoma, hypoglycaemia</td>
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<td></td>
<td>- Drugs (beta-receptor agonists, amphetamine, lithium, theophylline, caffeine, alcohol withdrawal)</td>
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<tr>
<td></td>
<td>- Essential tremor</td>
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<td></td>
<td>- Orthostatic tremor</td>
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<td></td>
<td>- Task-specific tremors (e.g. primary writing tremor)</td>
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<tr>
<td></td>
<td>- Parkinson’s disease</td>
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<td></td>
<td>- Corticobasal ganglionic degeneration</td>
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<td></td>
<td>- Dystonias</td>
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<tr>
<td>Peripheral neuropathies</td>
<td>- Charcot–Marie–Tooth syndrome</td>
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<td></td>
<td>- Paraproteinaemic neuropathies</td>
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<tr>
<td>Intention tremor</td>
<td>- Disease affecting cerebellum and its connections (e.g. dentate nuclei, superior cerebellar peduncles)</td>
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<td></td>
<td>- Multiple sclerosis</td>
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<tr>
<td></td>
<td>- Stroke</td>
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<td></td>
<td>- Tumour</td>
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<tr>
<td></td>
<td>- Drugs</td>
</tr>
<tr>
<td></td>
<td>- Cerebellar degenerations (e.g. paraneoplastic, spinocerebellar ataxias)</td>
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<td></td>
<td>- Rubral tremor</td>
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<tr>
<td>Hysterical tremor</td>
<td>- Mixed tremors not classifiable in any of the above</td>
</tr>
</tbody>
</table>

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**Figure T.28** Chest radiograph showing a left tension pneumothorax deviating the trachea to the right. This is a medical emergency.
TREMOR

RESTING TREMOR
The most important causes of resting tremor are Parkinson’s disease and other parkinsonian syndromes (e.g. multiple system atrophy). The cardinal features of Parkinson’s disease are slowness of voluntary movements (bradykinesia), rigidity and tremor. Typically, these symptoms are initially manifest as asymmetrical or unilateral limb clumsiness, often in one hand. Additional problems may develop, including postural instability, hypophonia (soft voice), dysarthria and facial hypomimia (poverty of expression). The tremor usually involves the heads, but may also involve the arms, legs, lower jaw or tongue. The tremor is coarse and regular, with a frequency of approximately 3–4 per second. The classical description is of pronation–supination of the forearm with flexion–extension of the fingers and adduction–abduction of the thumb, although this ‘pill-rolling’ is actually seen in only a minority of patients. The tremor can be exaggerated by distraction, for example by asking the patient to close their eyes and recite the days of the week. It may also be worsened by excitement or walking, where there may be reduced arm swing. Usually, the tremor itself causes little disability in daily tasks as it is markedly suppressed during volitional movements. The associated bradykinesia and rigidity, if present, do interfere with motor tasks. Rest tremor may be a feature of the parkinsonian variant of multiple system atrophy (MSA-P). Tremor is not characteristic of progressive supranuclear palsy, and is uncommon in drug-induced or post-encephalitis parkinsonism compared with idiopathic Parkinson’s disease.

POSTURAL TREMOR
The most important causes of a postural tremor are physiological tremor and essential tremor. Physiological tremor is a normal, small-amplitude oscillation that is usually asymptomatic, with a frequency of 8–13 per second. It most often affects the hands, and it is best elicited by asking the patient to hold their arms outstretched with the fingers wide apart. The tremor is not present when completely at rest. Many factors can exaggerate physiological tremor, including metabolic disturbances (hypothyroidism or hypeglycaemia) and intense anxiety or fright. Drugs including beta-receptor agonists, theophylline, caffeine and corticosteroids may increase physiological tremor. Essential tremor is the most common type of postural (or action) tremor. It is slower than physiological tremor, being of frequency 4–8 Hz, and usually affects the upper limbs. There may be an associated ‘No-No’ or ‘Yes-Yes’ head tremor that disappears when the head is supported. The jaw, lips, tongue and larynx can also be involved, causing voice disturbance, but isolated jaw or head tremor is reported not to be a feature of essential tremor. The disorder may be inherited in an autosomal dominant manner, with complete penetrance. Although it is often termed ‘benign essential tremor’, it may cause marked disability by preventing normal activities such as carrying objects accurately. It can be brought out during examination by asking the patient to maintain the arms outstretched. It is worsened by emotion, stress and fatigue, and may show a striking response to small doses of alcohol. Unlike Parkinson’s disease, the tremor is usually symmetrical, and asymmetrical onset (particularly in the legs) should call the diagnosis into question.

Orthostatic tremor may be considered a special form of postural tremor. Unlike essential tremor, it affects mainly the legs and is most prominent during standing. It diminishes with walking and disappears on sitting or lying flat. A buzzing noise may be heard on listening over the thigh or calf muscles with a stethoscope.

INTENTION TREMOR
The most important cause of intention tremor is cerebellar (and cerebellar outflow tract) disease, which may be due to many underlying disorders including demyelination, stroke, tumour or degenerative conditions. This tremor is perhaps inappropriately named as it occurs not on intention but on actual movement. Intention tremor can also be accurately termed ataxic, as it is always associated with cerebellar ataxia. Performing a precise guided movement, such as reaching with a finger to touch the examiner’s finger, brings it out. It is absent at rest or during the early part of a reaching movement, but is manifest as the body part approaches its target, requiring fine adjustments of position. It is irregular, of frequency 2–4 Hz, and continues for several cycles after the target has been reached. It may be disabling for fine motor task performance.

RUBRAL (HOLMES) TREMOR
This tremor is a mixture of resting, postural and intention tremor. Lifting the arm slightly, or maintaining an outstretched posture, causes a large-amplitude 2–5 Hz tremor that can be dramatic and cause the patient to lose balance. The lesion, as postulated by Gordon Holmes, an eminent British neurologist, was stated to be in the red nucleus of the midbrain (hence rubral). Experimental data now indicate that the tremor may be due to dentatothalamic cerebellar outflow fibres traversing the red nucleus. The most common causes of a rubral tremor are stroke, demyelination and Wilson’s disease.
Hysterical Tremor

Tremor is quite a rare manifestation of hysterical conversion. When it occurs, it is often confined to one limb, of large amplitude and is diminished by distraction – for example, performing a complex motor task with another body part. It is usually present equally at rest and during movement, unlike most organic tremors.

It should be remembered that tremors cannot always be fitted into the above groups, and that they may have features of more than one type. For example, in Parkinson's disease, a resting tremor may be combined with an intention tremor; in a cerebellar lesion, the tremor may (rarely) resemble a parkinsonian resting tremor.

Trismus (Lockjaw)

Mark Kinirons

(See Face, Abnormalities of Appearance and Movement, p. 176.)

Trismus, or lockjaw, signifies a maintained muscular spasm tending to closure of the jaws so that the mouth cannot be opened. The term does not include mechanical inability to open the jaws owing to such affections as mumps, alveolar abscess with surrounding inflammatory oedema, injury, Ludwig's angina, quinsy or severe tonsillitis, an odontoma, epiphlebitis of the mouth, myositis ossificans or cervicofacial actinomycosis. There are two conditions that may not at first sight be obvious but may lock the jaws together and simulate true trismus. These are impaction of a wisdom tooth and arthritic changes in the temporomandibular joint. Diagnosis is by careful local examination of the teeth and of the joint respectively; in the latter case, there may be arthritic changes in other joints as well. X-ray examination may be required to detect the joint changes or the impacted wisdom teeth.

Circumstantial evidence will generally serve to distinguish trismus due to hysteria or to facial neuralgia; any doubt at first experienced is dispelled if the patient is watched for a while. Convulsive seizures in a hysterical patient with trismus can generally be distinguished from those due to tetanus or to strychnine poisoning by their polymorphous character, and by the fact that touching the patient, and other similar stimulation, does not bring them on as certainly as would be the case with strychnine or tetanus.

In fits, for example epilepsy, the trismus is of short duration and offers no difficulty in diagnosis. Malingering may sometimes take the form of lockjaw, and it may be a little while before the fraud can be detected. In sleep, the malingering's muscles relax completely.

Trichiniasis is rare but, if infected pork is eaten raw or insufficiently cooked, the larvae of the parasites find their way to many different muscles, and they show a predilection for those of the tongue, mouth and jaws. The resultant irritation, pain and stiffness can cause trismus, the origin of which may be difficult to determine unless the history points to pork. The patient is very ill in the earlier stages, with high fever, and the condition may be fatal. The malady may be epidemic. The blood exhibits eosinophilia. The final criterion of the diagnosis is the discovery of the typical parasites coiled up in their little oval cysts among the affected muscle fibres.

Hydrophobia (rabies) and tetany seldom exhibit trismus as a prominent symptom. The former, although now almost unknown in Great Britain, would suggest itself if a convulsive illness developed after a bite by a dog, fox or other similar animal, particularly if the spasmodic muscular difficulty is markedly increased by efforts at swallowing. The symptoms may not develop for weeks or months after the bite, so that the patient may fall ill when he has come from a country overseas where rabies is endemic. Tetany, also rare, is at once distinguished by its typical carpopedal contractions; trismus, almost constant in tetanus, is nearly always absent in tetany.

Strychnine poisoning gives rise to generalized twitchings and convulsions long before trismus, the lateness of the development of the latter serving to distinguish it from tetanus. Furthermore, there is complete muscular relaxation between spasms. There may be evidence of strychnine having been taken or administered, either by mouth or hypodermically; the symptoms develop very acutely, and are often rapidly fatal.

Tetanus is the cause par excellence of trismus, which develops in days following introduction of the infection (e.g. from a small penetrating wound, burns, surgery, intravenous drug abuse or postpartum). Muscle stiffness starts usually in the neck muscles, spreading to those of the face and jaw, and thence to the rest of the trunk and limbs. Patients may have extremely painful generalized muscle spasms from the slightest stimulation (e.g. a feather stroke or the banging of a door). Muscle stiffness results in dysphagia, risus sardonicus and opisthotonos, and muscular relaxation is not possible unless an anaesthetic is given.

It may be possible to demonstrate the presence of the drumstick bacilli in films prepared from the deeper...
parts of the wound. Diagnostic difficulty may arise when there is no clear history, or when the wound has been so small that it has healed or cannot be found; even then, most cases are typical, particularly the combination of normal level of consciousness and normal cerebrospinal fluid despite recurrent generalized convulsions. Unnecessary anxiety in non-immunized individual may arise in cases of an impacted wisdom tooth, or of hysteria, where tetanus may at first be suspected; the subsequent course of the malady soon serves to exclude this. Involvement of the temporomandibular joint in a *serum reaction*, especially if prophylactic tetanus antitoxin has been given, may lead to the belief that tetanus has in fact set in.

Trismus may be simulated by *scleroderma* of the face. Here, however, the condition is rather one of fixation of the skin than of the muscles; the skin becomes like parchment so that one cannot pick it up between the fingers, it feels firm or almost hard, and the patient becomes unable to open the mouth properly. The disease is of slow onset and gradual progress, so that there is seldom difficulty in diagnosis.
UNDERACTIVITY

Mark Kiniron

The causes of underactivity are listed in Box U.1. Underactivity will most commonly be a normal response to fatigue and insomnia. The complaint of underactivity may also come from someone suffering an adjustment reaction from recent severe stress, such as marital or business problems or chronic illness. The response to stress may include anxious and depressed mood, physical complaints or disturbance of conduct including antisocial behaviour, or it may take the form of withdrawal from normal activities, without any clear evidence of mood change. Timely intervention with counselling may help the situation to resolve. Feelings of tiredness, fatigue and lack of drive accompany almost every form of debilitating illness, and a complaint of underactivity may be due, for example, to anaemia; endocrine disorders such as hypothyroidism or Addison's disease; cardiac, renal or hepatic impairment; chronic infections; or post-viral states.

Malnutrition caused by malabsorption will similarly cause symptoms related to debility. However, in people who are malnourished and also living in conditions of poverty and extreme deprivation, there can develop a profound state of apathy and underactivity as a result of a combination of adverse physical and social factors.

Many drugs of abuse are taken because they generate euphoria and underactivity. These include opioids, cannabis, hallucinogens (LSD, mescaline and magic mushrooms), barbiturates, benzodiazepines and other hypnotics and sedatives. More innocent drugs include codeine.

In depression, patients may show a slowing of all motor activity, including speech, gestures and facial movements. There is usually retardation of thinking, and a complaint of difficulty in initiating and executing all voluntary acts. Severe depression with retardation may pose diagnostic problems in an elderly person when a gradual onset may lead the clinician erroneously to diagnose dementia in a patient whose apparent memory loss and cognitive impairment are due to their slowness in performing the tasks rather than to real cognitive change. Pseudodementia due to depression often responds well to antidepressant treatments, and it is important to bear in mind that, during the early stages of treatment, retardation may improve more quickly than the mood state, and there is a high risk of a patient making a suicide attempt.

Depression is undoubtedly the most common cause of retardation, and even when retardation develops in a patient with another condition such as schizophrenia or obsessional illness, depression may be an important contributing factor. In obsessive-compulsive disorder (see OBSESSIONS, p. 470), some patients are so preoccupied by obsessional thoughts and inner rituals that they appear uncommunicative and slow in all their activities. In schizophrenia, underactivity may be the result of several different processes. Catatonic schizophrenia describes a relatively uncommon presentation of the psychosis, in which motor disorders dominate the picture. Patients may become mute or stuporous and, in rare cases, adopt strange postures, exhibiting a disturbance of muscle tone by which their limbs will remain in any new position – however uncomfortable – for minutes at a time (waxy flexibility). They may show automatic obedience, responding to all requests without question; or negativism, responding in exactly the opposite way to that requested; or they will sometimes remain mute and unresponsive, unwilling to engage at all with the interviewer. A detailed history and follow-up assessment is the only way that the diagnosis can be confirmed.

A rather more common type of underactivity affecting many patients with schizophrenia occurs with so-called ‘negative’ symptoms. These patients lose their sense of drive and volition, and they become apathetic and careless about their appearance. They have a paucity of speech and emotional blunting that limits their full enjoyment of anything. They are poor time-keepers at work, and will be described as lazy, egotistical and inconsiderate by family and friends who are not made aware of the true nature of the condition. Depressive symptoms are common in chronic schizophrenia, and these may exacerbate the underlying slowness.
and apathy. The side effects of antipsychotic medication are a further cause of slowness in these patients, since higher doses of most antipsychotic drugs cause a combination of motor and psychological slowing, which adds to the negative symptoms.

_S stupor describes a state in which a subject is fully conscious but makes no spontaneous movements and does not respond to stimuli. Occasionally, this may be psychogenic in origin, in which case the onset is sudden and stress-related, and the patient may sit motionless for long periods without moving or talking, although their muscle tone, posture and eye movements indicate that they are neither asleep nor unconscious._

Patients with damage to their frontal lobes show changes in behaviour, mood and volition, typically with loss of initiative and spontaneity and a marked reduction in motor behaviour. Such people have particular difficulties in starting any new initiatives. Despite their mood—which can be euphoric—they perform tasks slowly and incompetently. A frontal lobe syndrome may follow head injury or may be due to vascular or space-occupying lesions, or other pathologies such as demyelination involving the frontal lobes.

**UPPER LIMB, PAIN IN**

Fred Heatley & Jonathan Compson

**PRINCIPLES OF DIAGNOSIS**

The presenting features of disease and injury of the upper limb are pain, deformity, stiffness, weakness and paraesthesiae, swelling, instability and loss of function. Pain is by far the most common symptom. This may be localized (e.g. the pain of De Quervain’s tenosynovitis, which localizes to the radial styloid) or, more often, presents as a pattern. Dual pathology is common, for example cervical spondylosis and carpal tunnel syndrome. Routine examination therefore involves examination of the whole arm, including the cervical spine, the neurology and the vascular system. In comparison to the lower limb, nerve entrapments and tendinitis/tenosynovitis are more common, while vascular disease is less frequent.

**PAIN REFERRED INTO THE ARM**

This falls into two main categories. Sharp, well-localized neuralgia often associated with paraesthesia is usually caused by nerve root or trunk compression. Diffuse discomfort in the upper limb, which is often difficult for the patient to describe and which may be accompanied by changes in skin temperature, vascularity and sweating, suggests involvement of the autonomic pathways. With this type of ‘cylindrical’ limb pain, an origin within the thorax or the thoracic spine, or involvement of T1 nerve root/stellate ganglion, should be sought. Many musculoskeletal pains are also transmitted via the autonomic system. For example, the pain from a frozen shoulder can temporarily be relieved by blocking the stellate ganglion with local anaesthetic.

**LESIONS IN THE CERVICAL SPINE (BOX U.2)**

It should be noted that X-ray changes of cervical spondylosis are a normal finding after the age of 40 years. Over the age of 60, neurological symptoms and signs referred from the cervical roots are common. Therefore, great care must be taken before ascribing patients’ symptoms solely to spondylosis, as there is often dual pathology.

_Cervical spondylosis can produce three clinical syndromes, which may occur alone or in combination: (i) pain and stiffness of the neck, which is often recurrent and may be aggravated by tension, anxiety and posture; (ii) radicular pain radiating down one or both arms, and which may or may not be associated with muscle wasting, weakness and reflex changes (this often being referred to as brachial neuralgia); and (iii) compression of the cervical cord, which may produce three sets of symptoms and signs:_

- Weakness, wasting and fibrillation in the upper limbs, with reduction or loss of the tendon reflexes at the level of the compression.
Paraesthesiae in the arms and legs, with or without impaired sensation in the hands and feet
Pyramidal tract involvement, with weakness, spasticity, hyperreflexia and extensor plantar responses in the feet

The combination of weakness and wasting in the arms and spastic weakness in the legs resembles amyotrophic lateral sclerosis; spondylosis may usually be distinguished from this by the history of paraesthesiae, evidence of sensory impairment and radiographic or magnetic resonance imaging (MRI) showing cord compression. L'Hermitte’s sign may be demonstrable – an electric shock sensation on neck flexion.

Disc herniation at the C5/C6 and C6/C7 intervertebral spaces is a common cause of pain in the upper limb. The onset may be acute, with well-localized pain radiating from the back of the neck, across the back of the shoulder, and down the arm and forearm to the wrist or fingers. More commonly, the onset is less dramatic, often after a period of recurrent aching and stiffness in the neck. Pain may be aggravated by movements of the neck, by downward pressure on the head (the compression test) and by changing the position of the arm. Pain relief by applying traction implies entrapment at an exit foramen – with the patient sitting, place one hand under the mandible and the other under the occiput, and lift. Pain may radiate downwards into the scapular region and to the upper chest. Sensory disturbances are common, and they may be detected in a dermatomal distribution (Fig. U.1). Muscle weakness may be detected in the appropriate muscles. The clinical signs associated with the most common root lesions are indicated in Table U.1. Depression of the biceps jerk indicates a lesion of the C5 root; paraesthesiae in the thumb and index finger with depression of the brachioradialis jerk indicates a lesion of the C6 root; and paraesthesiae in the index and middle fingers with loss of the triceps jerk are associated with a lesion of the C7 root. Note that there are no specific reflexes for the C8 or T1 roots. For C8, test the power of finger flexion, and check sensation in the little finger and the ulnar border of the hand. For T1, evaluate the intrinsic muscles of the hand, in particular the power of finger abduction. Ask the patient to ‘spread out’ the fingers; the examiner then squeezes them together. Compare both hands, and test the sensation of the ulnar border of the elbow and the upper arm. Check for paraesthesiae in the feet and spasticity in the legs, and examine the plantar response in case there is associated cord compression.

X-rays of the cervical spine may show disc space narrowing, especially at the C5/C6 or C6/C7 level, with lipping of the adjacent margins of the vertebral bodies. In the acute stage, X-rays may not reveal a relevant abnormality, as disc space narrowing in the lower cervical spine is an extremely common appearance in normal individuals over the age of 40. Computed tomography (CT) or magnetic resonance imaging (MRI) is the investigation of choice in demonstrating a disc herniation, including herniation into the lateral recess. Spinal fluid examination is not

**Table U.1 Signs and symptoms associated with common nerve root lesions affecting the arms**

<table>
<thead>
<tr>
<th>Root</th>
<th>Paraesthesiae/numbness</th>
<th>Muscle weakness</th>
<th>Reflex change</th>
</tr>
</thead>
<tbody>
<tr>
<td>C5</td>
<td>Radial aspect of arm</td>
<td>Shoulder abduction</td>
<td>Biceps jerk diminished</td>
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<tr>
<td></td>
<td></td>
<td>Elbow flexion</td>
<td></td>
</tr>
<tr>
<td>C6</td>
<td>Thumb and index finger</td>
<td>Wrist extension and pronation</td>
<td>Brachioradialis jerk diminished</td>
</tr>
<tr>
<td>C7</td>
<td>Middle finger, back of hand</td>
<td>Elbow extension and finger extension</td>
<td>Triceps jerk diminished</td>
</tr>
<tr>
<td>C8</td>
<td>Little finger, ulnar border of hand</td>
<td>Finger and wrist flexion</td>
<td>–</td>
</tr>
<tr>
<td>T1</td>
<td>Ulnar border of forearm and elbow</td>
<td>Intrinsic muscles of hand</td>
<td>–</td>
</tr>
</tbody>
</table>

N.B.
1 The biceps jerk is predominantly C5, but also has a component from C6.
2 The brachioradialis jerk, C6, is often, although erroneously, called the supinator jerk.

**Figure U.1** Dermatomes of the arm from C4–8 and T1–2.
a routine investigation for disc prolapse as it is usually normal; however, it is worth recalling that the protein content may be raised in a large herniation, especially in the presence of cord compression.

Other causes of brachial neuralgia are uncommon, but viral, bacterial and fungal infections not only occur but can also be diagnostically deceptive in their early stages. Herpes zoster is a good example. Later, the presence of a vesicular rash and residual pigmented scars in a dermatomal distribution provide an obvious explanation for the persistent pain. Weakness of one or more muscles in the limb with cutaneous hyperalgesia or hypoesthesia may occasionally be present. Acute viral infections (acute viral radiculitis) can affect individual components of the brachial plexus causing pain, which can be severe, weakness or paralysis in a similar manner to poliomyelitis. The classic example is winging of the scapula due to involvement of the long thoracic nerve resulting in paralysis of serratus anterior. In modern practice, this is more frequently caused by trauma since, due to the anatomy, this nerve is vulnerable to compression (e.g. backpacker’s shoulder) or repetitive minor trauma (e.g. weight lifting) (see Fig. U.6).

Vertebral and paravertebral abscesses may result from tuberculosis or brucellosis or be caused by more common pyogenic organisms such as Staphylococcus aureus. The incidence of spinal infection is increasing due to the rise in drug addiction and AIDS. Fungal or parasitic lesions are occasionally encountered in both these groups. Such lesions may or may not be accompanied by fever, and initial symptoms may closely resemble a cervical disc prolapse. Occasionally, there are no other pointers to a septic lesion, so that severe root symptoms in the arms together with paraesthesiae, widespread atrophy, loss of reflexes and variable sensory loss; more than one root is implicated. Positive syphilitic serology alone should not be taken to indicate this rare condition in the absence of other diagnostic features.

Primary (rare) or secondary neoplasms of the vertebral bodies may give rise to root pain with or without motor, sensory and reflex changes. X-rays are usually abnormal – collapse of a body, loss of a pedicle, and cystic or sclerotic lesions are the common features, although only approximately 50 per cent of metastases will show on plain films. In multiple myeloma, X-rays may show nothing more than diffuse osteoporosis until the vertebral body collapses. Technetium bone scanning can be helpful, although spurious ‘hot spots’ may be seen in the presence of marked degenerative disease of the spine, and it is important to bear in mind that plasmacytomas and myeloma deposits may (as with plain X-rays) not be detected by this technique. CT and MRI scanning are particularly helpful in early lesions and also in determining the extent of any extra-osseous extension. In 60 per cent of cases of multiple myeloma, Bence Jones protein (which precipitates at 60 °C) is present in the urine. Electrophoresis of the serum shows a distinctive spike close to the gamma position. Tumours of the meninges and roots usually cause symptoms in the legs from compression of the pyramidal and sensory tracts, as well as pain in the arm. Scalloping of the pedicles is the classical picture on plain X-rays. Root lesions in the presence of multiple cutaneous neurofibromata (von Recklinghausen’s disease) should raise the possibility of transformation to a malignant neurofibrosarcoma. Syringomyelia occasionally causes pain in the arm, but only as a late feature. By this stage, the classical features of dissociated sensory loss, muscle wasting and hyporeflexia in the arms with pyramidal signs below the level of the lesion are likely to be apparent.

Traumatic injuries to an otherwise intact cervical spine are remarkably easy to miss. There are two reasons for this. First, these injuries are not all due to high-velocity forces; for example, a ‘hangman’s fracture’ – a fracture through the pedicles of C2 – can occur with a blow to the forehead in a relatively straightforward fall while skiing. The second reason for misdiagnosis is that X-rays, particularly of the cervicothoracic junction, are difficult to interpret due to the overlying shadows of the shoulders on the lateral views. Any patient presenting with stiffness and spasm in the cervical spine, especially if there is a radiating pain into an arm, should be presumed to have major cervical spine injury until proven otherwise, and not assumed merely to have a whiplash injury. Fracture dislocations of the cervical spine are especially likely in the presence of rheumatoid arthritis or ankylosing spondylitis. In the former condition, atlanto-axial and/or subaxial subluxation of the spine may lead
to upper and lower limb symptoms (see NECK, PAIN AND/OR STIFFNESS, p. 450; and Figs N.17–N.19, p. 453–5), while the fused segments of a spondylitic spine are particularly at risk of fracture with or without displacement. Fractures of cervical vertebrae due to osteoporosis are unusual.

**LESIONS OF THE BRACHIAL PLEXUS AND SUBCLAVIAN/BRACHIAL ARTERIES**

The causes of these lesions are as follows:

- **Thoracic outlet syndrome**
  - Cervical rib/fibrous band
  - Scalenus anterior syndrome
  - Subclavian aneurysm
  - ‘Drooping shoulder’ syndrome
- **Malignant infiltration**
  - axillary lymph nodes (e.g. lymphoma, metastatic carcinoma)
  - posterior triangle disease (e.g. metastatic carcinoma)
  - apical bronchial carcinoma (Pancoast tumour)

The *thoracic outlet syndrome* is due to compression of the neurovascular bundle. Causes include an abnormal insertion of the scalenus anterior muscle, the presence of a cervical rib or a vestigial fibrous band, and poor posture from drooping of the shoulder with consequent stretching of the plexus over a normal first rib (most commonly seen in middle-aged overweight women). Typically, pain is felt behind the clavicle and down the inner aspect of the arm. Paraesthesiae and hypoaesthesia in the C8 and T1 dermatomes, with associated vasospastic features, are common findings (i.e. in the little finger and ulnar border of the hand and forearm). Symptoms may be aggravated by carrying heavy weights, although this is not diagnostic. There may be atrophy of the hypothenar eminence and interosseus muscles. The diagnosis is usually based on an induction of paraesthesiae and numbness by abduction of the arm to 90° with external rotation and detection of an arterial bruit in the supraclavicular fossa during this manoeuvre with disappearance of the symptoms and bruit on returning the arm to the neutral position. Finding a position of the arm in which the radial pulse is obliterated has been considered a key diagnostic finding. However, this may be demonstrated in normal subjects, and symptoms can be due to compression of the brachial plexus without involvement of the subclavian artery. The diagnosis is not, therefore, dependent on a demonstration of arterial compression. When chronic or recurrent subclavian artery compression is present, this may (occasionally) lead to the development of aneurysmal dilatation of the subclavian artery, and consequently to emboli causing infarcts in the fingers. In a few patients, the accessory rib may be palpable and also visible on X-ray; not infrequently, the rib is vestigial, being replaced by a fibrous band that cannot be detected (see Fig. M.15). There is a tendency to overdiagnose thoracic outlet syndrome as a cause of pain in the arm. The alternative diagnoses of cervical spondylosis, cervical disc lesion or a peripheral nerve lesion are much more common.

Pain in the arm is occasionally due to pressure on, or infiltration of, the brachial plexus by *malignant tumours*. Lymphadenopathy associated with lymphomas or metastatic carcinoma will usually be detectable by palpation of the axilla and of the posterior triangle of the neck. However, infiltration of the plexus by metastatic carcinoma, especially from the breast, may take time to become evident. Involvement of the plexus by upward spread of an apical bronchial carcinoma (Pancoast tumour), or more rarely by apical inflammatory lung disease, can produce unilateral Horner’s syndrome in addition to arm pain. Such lesions can usually be detected on a chest X-ray (Fig. U.2). In all of these conditions, severe pain may be present without any accompanying signs in the early stages. Further infiltration usually leads to paralysis, with relative sparing of sensation.

![Figure U.2 Chest radiograph of a patient with Horner's syndrome, showing a left Pancoast lung tumour (arrowed), a classic cause of referred arm pain, weakness of the small muscles of the hand (T1 innervation) and unilateral Horner's syndrome.](image-url)
LESIONS IN THE THORAX, THORACIC SPINE AND ABDOMEN

The causes of these lesions are as follows:

- Cardiac ischaemia
- Syphilitic aortitis
- Thoracic disc
- Oesophagitis
- Diaphragmatic irritation
  - Subphrenic abscess
  - Ruptured abdominal viscus
  - Lesions of the spleen and pancreas

In contrast to the characteristically searing localized pain of nerve root involvement, pain in the arm originating in the chest has a dull, poorly localized quality, sometimes described as cylindrical. Autonomic controlled functions, including temperature of the arm and sweating, are often altered.

Pain associated with *myocardial infarction* and the exercise or stress-related pain of angina pectoris is usually readily recognized and confirmed by electrocardiogram (ECG) or exercise testing.

*Syphilitic aortitis* may induce similar referred pain. *Oesophagitis* can produce cylindrical arm pain with or without more classical ‘heartburn’. Such pain may also be accompanied by ECG abnormalities, so that accurate distinction from myocardial ischaemia may rest upon exercise testing, a trial of glyceryl trinitrate and visualization of the upper gastrointestinal tract. Referral upon exercise testing, a trial of glyceryl trinitrate and distinction from myocardial ischaemia may rest without more classical ‘heartburn’. Such pain may also induce similar referred pain.

Pain of nerve root involvement, pain in the arm originating in the chest has a dull, poorly localized quality, sometimes described as cylindrical. Autonomic controlled functions, including temperature of the arm and sweating, are often altered.

Pain associated with *myocardial infarction* and the exercise or stress-related pain of angina pectoris is usually readily recognized and confirmed by electrocardiogram (ECG) or exercise testing.

*Syphilitic aortitis* may induce similar referred pain. *Oesophagitis* can produce cylindrical arm pain with or without more classical ‘heartburn’. Such pain may also be accompanied by ECG abnormalities, so that accurate distinction from myocardial ischaemia may rest upon exercise testing, a trial of glyceryl trinitrate and visualization of the upper gastrointestinal tract. Referral from the thoracic spine is a little recognized cause of aching in the arm or ‘fibrositis’. *Thoracic disc prolapse* usually presents with an insidious onset of thoracic back pain, which is occasionally referred into the arm. There is local thoracic spine tenderness, best elicited by sequentially pressing on the thoracic vertebrae with the patient prone. Pain is exacerbated by thoracic rotation. Other causes of stiffness of the thoracic spine, including spondylosis, may lead to similar symptoms.

In a minority of instances, myocardial infarction leads to the development of pain and stiffness at one shoulder with varying degrees of pain, swelling, osteoporosis, vasomotor disturbance and trophic skin changes more distally in the limb. This ‘*shoulder–hand syndrome*’ is an example of a reflex sympathetic dystrophy or algodystrophy. This can also occur following nerve injuries, fractures such as a Colles fracture or a sprained wrist. The pain is usually described as ‘burning’. The fingers, wrist and shoulder stiffen, but the elbow is less affected. The condition can last for many months. Local anaesthetic blockade of the stellate ganglion is often effective in relieving the pain, thereby confirming the diagnosis and shortening the period of disability. However, when severe, full finger mobility in particular does not usually recover.

Since the C5 nerve root supplies the diaphragm and the shoulder, several upper abdominal conditions can present with shoulder pain. Examples include subphrenic abscess, lesions of the spleen and pancreas, and subdiaphragmatic irritation from a perforated viscus.

PAIN, STIFFNESS AND WEAKNESS OF THE SHOULDER GIRDLE

This includes the clavicle, acromioclavicular joint, scapula and glenohumeral joint, which is considered in the next section. A good knowledge of the anatomy is essential to making a diagnosis (Box U.3).

Pain may be referred, may arise locally or may be a combination of both – for example, cervical spondylitis and a frozen shoulder often co-exist. Pain situated over the acromioclavicular joint usually indicates pathology within the joint. Glenohumeral pain is, classically, referred over the deltoid on the lateral aspect of the proximal humerus, the area supplied by the C5 root. Glenohumeral osteoarthritis often presents with a posterolateral pain, while pain from bicipital tendinitis is felt anteriorly over the bicipital groove. The most common cause of stiffness is a frozen shoulder. Other causes include: glenohumeral osteoarthritis, a condition that used to be relatively rare but that has become much

<table>
<thead>
<tr>
<th>Box U.3 Shoulder girdle disorders excluding the glenohumeral joint</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Congenital</strong></td>
</tr>
<tr>
<td>- Unilateral elevation of the scapula (Sprengel’s shoulder)</td>
</tr>
<tr>
<td>- Bilateral elevation of the scapula (Klippel–Feil syndrome)</td>
</tr>
<tr>
<td>- Absence of the clavicle (craniocleidodyostosis)</td>
</tr>
<tr>
<td><strong>Traumatic</strong></td>
</tr>
<tr>
<td>- Sternoclavicular joint dislocation/subluxation</td>
</tr>
<tr>
<td>- Anterior</td>
</tr>
<tr>
<td>- Posterior</td>
</tr>
<tr>
<td>- Spontaneous</td>
</tr>
<tr>
<td>- Clavicular fractures</td>
</tr>
<tr>
<td>- Acromioclavicular dislocation/subluxation</td>
</tr>
<tr>
<td><strong>Infective</strong></td>
</tr>
<tr>
<td>- Osteomyelitis of clavicle</td>
</tr>
<tr>
<td>- Chronic recurrent multifocal osteomyelitis – the clavicle is one of the sites involved</td>
</tr>
<tr>
<td>- Septic arthritis of the sternoclavicular joint</td>
</tr>
<tr>
<td><strong>Septic arthritis of the acromioclavicular joint</strong></td>
</tr>
<tr>
<td>- Tuberculous synovitis of the acromioclavicular joint</td>
</tr>
<tr>
<td><strong>Inflammatory</strong></td>
</tr>
<tr>
<td>- Rheumatoid arthritis</td>
</tr>
<tr>
<td><strong>Degenerative</strong></td>
</tr>
<tr>
<td>- Osteoarthritis of the acromioclavicular joint</td>
</tr>
<tr>
<td><strong>Miscellaneous</strong></td>
</tr>
<tr>
<td>- Osteonecrosis of the medial clavicular epiphysis (Friedreich’s)</td>
</tr>
<tr>
<td>- Condensans osteitis of clavicle</td>
</tr>
<tr>
<td>- Paralysis of serratus anterior (‘Winging’ of the scapula - see p. 705)</td>
</tr>
<tr>
<td>- Traumatic</td>
</tr>
<tr>
<td>- Post-viral neuritis</td>
</tr>
<tr>
<td>- Muscular dystrophies (e.g. Duchenne’s)</td>
</tr>
</tbody>
</table>
more frequent due to the longevity of the population; and poorly treated trauma, for example a missed posterior dislocation of the shoulder. Rotator cuff tears are the common cause of weakness, but these need to be distinguished from other causes of deltoid weakness, especially a C5 neuropathy, a lesion of the axillary nerve, or neurological diseases such as motor neurone disease. A combination of the age of the patient plus the main symptom often gives a good indication of the diagnosis. An elderly patient with pain, weakness and stiffness is likely to have osteoarthritis, a middle-aged patient with a painful stiff shoulder probably has a frozen shoulder, and a young man with pain and reduced rotation is likely to have an unstable shoulder.

Do not examine the shoulder in isolation. The whole of both upper limbs must be visible and examined from the back as well as the front. Generalized diseases also affect the shoulder, although most (e.g. rheumatoid arthritis) are usually obvious. Look for deformities – a prominence over the distal end of the clavicle probably indicates acromioclavicular disease; a high-riding scapula is a Klippel–Feil or Sprengel shoulder. Wasting of the deltoid occurs in most shoulder conditions but, if excessive, check particularly for a C5 palsy or an axillary nerve paralysis with the loss of sensation over the lateral aspect of the shoulder. Localizing any tenderness is very helpful, so palpation should follow a routine – the sternoclavicular joint, the clavicle, the acromioclavicular joint and the acromion, moving along its sharp border from front to back. With the shoulder extended, the biceps tendon lies just in front of the acromion; rotate the humerus to bring the prominence adjacent to the bicipital groove into play under your finger. Move across the front of the humeral head to palpate the coracoid process lying in the deltoid pectoral groove. Palpate around the scapula, moving posteriorly onto the thoracic and then the cervical spine. The shoulder girdle has an enormous range of movement, but for normal function there must be normal movements in three joints – sternoclavicular, acromioclavicular and glenohumeral – plus full scapulothoracic movement. The split between glenohumeral and scapulothoracic movement is approximately 2 to 1. Again, compare both shoulders from the front and the back. A useful ‘ready reckoner’ for normality is the Apley ‘opposite scapula three scratch test’ by feeling the opposite scapula from three directions – arm across the chest, arm behind the neck, and arm behind the back. Shoulder movements can be very deceptive since a disease affecting the glenohumeral joint will not abolish scapulothoracic movement; for example, a frozen shoulder in which there is minimal glenohumeral movement may still have 90° of abduction due purely to scapulothoracic movement. When looking for impingement from rotator cuff lesions, abduct both arms simultaneously. Rotator cuff impingement pain occurs in the arc from 60 to 120°, while pain in the arc above 120° is likely to arise from the acromioclavicular joint. Finally, test the passive movements to determine whether or not they match or exceed the active range. Surprisingly, the parents of a child with Sprengel’s shoulder usually complain that the normal scapula is too low rather than the abnormal scapula is too high. Shoulder abduction may be somewhat restricted but, in most cases, function is satisfactory. Occasionally, a Sprengel shoulder can be bilateral, and there may be associated deformities such as scoliosis. In Klippel–Feil syndrome, failure of shoulder descent is bilateral and is associated with abnormalities of the cervical spine. The neck seems absent, and the patient has the appearance of wearing a clothes-hanger permanently under the jacket (Fig. U.3). In craniocleidodysostosis, the clavicles are absent. This permits a remarkable amount of movement, so that both shoulders can actually be pressed together in front of the thorax (Fig. U.4).
A number of other abnormalities can be present, including an altered shape of the skull and infantile coxa vara of the hips. Osteomyelitis of the clavicle and septic arthritis of the acromioclavicular joint are usually obvious as the bone and joint are superficial. However, infections of the sternoclavicular joint can be deceptive, and are a cause of ‘pyrexia of unknown origin’ (PUO) in patients in intensive care units, those who are immunocompromised and drug addicts. Spontaneous dislocation of the sternoclavicular joint affects children and young adults, the clavicle dislocating when the arm is elevated above the head. It is often bilateral, rarely causes pain and is best left alone. Traumatic dislocation results from either a direct force or an indirect compressive force with the shoulders being squeezed together. In the early stages, a haematoma often masks the physical signs. Posterior dislocations are fortunately much rarer than anterior dislocations as they are associated with major complications to the many important structures lying immediately posterior. Clavicular fractures are common. The usual mechanism is a fall on the point of the shoulder (and not on the outstretched hand, as stated in many older books). It is also the most common birth injury fracture. Most fractures (approximately 75 per cent) affect the middle third. These fractures unite readily, and a non-union is a rare complication. Fractures of the lateral third, approximately 20 per cent, can be more troublesome. They can occur across the interval between the attachment of the conoid and trapezoid ligaments, extend into the acromioclavicular joint or in children cause a pseudo-disarticulation in which the distal clavicle is avulsed out of its periosteal sleeve. Injuries to the acromioclavicular joint are common (Fig. U.5). Unlike the majority of synovial joints, the main ligaments are not a thickening of the capsule; they are situated proximally and run from the coracoid process to the underside of the distal clavicle as the conoid and the trapezoid ligaments. The cause is a fall onto the tip of the shoulder, the acromion and consequently the scapula, together with its coracoid process being depressed downwards and away from the clavicle. Injuries vary from a sprain of the capsule, through partial and complete tears of the conoid/trapezoid ligament complex, to gross disruption of the aponeurosis of the overlying deltoid and trapezius muscles, the latter injuries being due to high-velocity force. The minor sprains result in tenderness around the joint; with disruption of the capsule, there is subluxation and more pronounced pain and swelling; while rupture of the conoid/trapezoid ligament complex will permit dislocation of the clavicle from the acromion, causing an obvious deformity – best visualized from behind with the patient sitting and the arm hanging down by the side. These injuries are frequently missed or minimized on routine shoulder X-rays. The correct view is an anteroposterior film taken with a 15° superior tilt and the arm unsupported. Rheumatoid arthritis of the acromioclavicular and sternoclavicular joints may affect up to 60 per cent of patients with rheumatoid disease. This is frequently overlooked, but it is a potent cause of unnecessary loss of shoulder function as it is easily treated. Pain is usually felt anteriorly and radiates across the point of the shoulder. Demonstrating the degree of destruction requires a 15° tilted anteroposterior X-ray. Patients often present late, with fixed internal rotation and very little other movement remaining in the shoulder. Osteoarthritis is an important entity since osteophytes on the underside of the joint are a major cause of subacromial impingement and tears of the rotator cuff. The pain is usually well localized to the joint and is exacerbated by using the arm above the head. The joint is swollen and tender, and abduction produces a high painful arc (i.e. above 120°). The diagnosis may be confirmed by injecting local anaesthetic, which abolishes the painful arc. Osteonecrosis of the medial clavicular epiphysis (Friedreich's disease) is a rare condition that is usually only diagnosed in retrospect by the late development of osteoarthritis of the joint. Condensans osteitis also affects the medial end of the clavicle, but the joint space is preserved. It is usually seen in women in the middle years. They present with pain at the medial end of the clavicle, and X-rays confirm the diagnosis.

Figure U.5 Prominence of the distal end of the clavicle due to dislocation at the acromioclavicular joint (arrowed).
Winging of the scapula usually results from trauma to the long thoracic nerve (C5, C6 and C7), which supplies serratus anterior – ‘back-packer’s shoulder’ (Fig. U.6). A spontaneous onset is attributed to a post-viral neuritis.

Paralysis of the trapezius muscle is due to injury to the spinal accessory nerve, which is easily damaged by operations, including ‘minor’ operations, to remove a lump in the neck. Loss of trapezius results in drooping of the shoulder and an inability to control scapulothoracic movement, thus causing a major loss of shoulder function.

**INJURIES AND DISEASES OF THE GLENOHUMERAL JOINT**

Testing the shoulder muscles should follow the procedure in Table U.2.

See Box U.4 for a summary of the diagnostic categories for pain arising from the glenohumeral joint and humerus.

**Lesions of the rotator cuff**

The rotator cuff is a common source of symptoms as it is liable to: (i) compression between the humeral head and the acromion/coracoacromial ligament; and (ii) degenerative tears due to its relatively poor blood supply. Osteophytic lipping on the inferior aspect of the acromioclavicular joint or the anterior lip of the acromion may further reduce the space for the supraspinatus component of the cuff, and consequently this is the most common site for impingement. The classical presentation is the ‘painful arc’ which typically occurs between 70 and 120° of active abduction. The pain is usually localized to the anterolateral aspect of the shoulder, with radiation down to the deltoid insertion. It is a common condition, particularly in older athletes and in those whose work puts repetitive demands on the shoulder, for example carpenters. The syndrome is relatively uncommon under the age of 40.

In addition to the standard examination of the shoulder and the demonstration of a painful arc in

**Box U.4 Diagnostic categories for pain arising from the glenohumeral joint and humerus**

<table>
<thead>
<tr>
<th>Lesions of the rotator cuff</th>
<th>Infection</th>
</tr>
</thead>
<tbody>
<tr>
<td>Impingement syndromes</td>
<td>Acute pyogenic</td>
</tr>
<tr>
<td>Cuff rupture</td>
<td>Septic arthritis</td>
</tr>
<tr>
<td>Acute calcification</td>
<td>Osteomyelitis</td>
</tr>
<tr>
<td>Frozen shoulder</td>
<td>Chronic</td>
</tr>
<tr>
<td>Instability</td>
<td>– Tuberculosis</td>
</tr>
<tr>
<td>Anterior subluxation/dislocation</td>
<td>– Infected implant surgery</td>
</tr>
<tr>
<td>Posterior subluxation/dislocation</td>
<td>Arthritis</td>
</tr>
<tr>
<td>Multidirectional instability</td>
<td>Rheumatoid arthritis</td>
</tr>
<tr>
<td></td>
<td>Osteoarthritis</td>
</tr>
<tr>
<td></td>
<td>– Primary</td>
</tr>
<tr>
<td></td>
<td>– Post-traumatic</td>
</tr>
<tr>
<td></td>
<td>– Post-rotator cuff disintegration</td>
</tr>
<tr>
<td></td>
<td>Osteonecrosis</td>
</tr>
<tr>
<td></td>
<td>– Post-traumatic</td>
</tr>
<tr>
<td></td>
<td>– Steroids, alcohol, etc.</td>
</tr>
<tr>
<td></td>
<td>– Haemoglobinopathies (e.g. sickle-cell disease)</td>
</tr>
<tr>
<td></td>
<td>– Caisson’s disease (the ‘bend’s)</td>
</tr>
</tbody>
</table>

**Table U.2 Put the muscle under test in the optimal position and test against resistance**

<table>
<thead>
<tr>
<th>Muscle</th>
<th>Position</th>
</tr>
</thead>
<tbody>
<tr>
<td>Deltoid</td>
<td>Test all three components (i.e. flexion, abduction and extension)</td>
</tr>
<tr>
<td>Supraspinatus</td>
<td>In plane of scapula (i.e. slight flexion and 15° of abduction)</td>
</tr>
<tr>
<td>Subscapularis</td>
<td>Internal rotation against resistance with the elbow flexed to 90° and held behind the back (get the patient to lift the hand away from the back/buttocks)</td>
</tr>
<tr>
<td>Infraspinatus</td>
<td>External rotation against resistance with the elbow flexed to 90° and held by the side</td>
</tr>
<tr>
<td>Trapezius</td>
<td>Shrug shoulders upwards</td>
</tr>
<tr>
<td>Serratus anterior</td>
<td>Press with both hands against a wall. Weakness causes ‘winging’ of the scapula (see Fig. U.6)</td>
</tr>
</tbody>
</table>

**Figure U.6** Winging of right scapula secondary to paralysis of serratus anterior. The scapula becomes prominent when the patient pushes against a wall with his hands.
abduction, there are two useful special tests: (i) the impingement test – press down on the shoulder with one hand and forcibly fully flex the arm with the other hand; (ii) place the arm in forward flexion at 90° with the elbow at 90° and then passively internally rotate the humerus. An injection of local anaesthetic into the subacromial space should abolish the pain of impingement.

The subacromial space is best visualized by a lateral X-ray taken along the line of the scapula with a 5–10° caudal tilt. This will reveal any calcification in the tendon or abnormalities of the acromion. An anteroposterior view with a 15° superior tilt will show the acromioclavicular joint. CT and MRI imaging are excellent for showing the rotator cuff (Fig. U.7), as is ultrasound examination by an experienced doctor or radiographer.

**Acute calcification** most commonly affects the supraspinatus tendon, followed by the infraspinatus and teres minor. Subscapularis calcification is uncommon. It is a disease of middle age, usually between 40 and 50 years, and is rarely seen over 70 years. While the majority of patients are asymptomatic, the calcification showing as an incidental finding on plain X-rays, a few present with severe pain and inability to move the arm (Fig. U.8a). As the calcification disperses (Fig. U.8b) the pain settles, usually over 5–7 days and the shoulder returns to normal, although a painful arc may persist.

Ruptures of the rotator cuff (Fig. U.7) begin on the undersurface of supraspinatus tendon near its insertion, progressing to a full-thickness tear and then spreading posteriorly to involve the infraspinatus. The rotator cuff retracts posteriorly and inferiorly, allowing the humeral head to migrate through the gap, so that it wears against the acromion. The onset can be acute or gradual, and it is often precipitated by relatively minor trauma, for example, grabbing something to prevent a fall. In addition to the features of impingement, which are commonly present, there is evidence of weakness. The **frozen shoulder** is a painful, stiff shoulder secondary to a global loss of glenohumeral movement. It is more common in women than men, with an average age on presentation of 55 years (range 40–70 years). It is of slow onset and has three clinical stages, each of which lasts between 3 and 9 months – the painful stage, the painful stiff stage and the resolving stage. It rarely recurs in the same shoulder, but it can affect the opposite shoulder.
In the vast majority of cases, the shoulder returns to near-normality. Characteristically, there is slow onset of painful restriction of movement, especially external rotation and abduction, with the pain being most prominent at night. Biopsies show an active fibroblast proliferation and transformation to myofibroblasts in the capsule and the coracohumeral ligament. There is an increased incidence in patients with diabetes, hyperthyroidism and Dupuytren’s contracture. Radiologically, the joint is normal apart from some osteopenia. This, together with the history, will distinguish a frozen shoulder from other causes of a stiff painful shoulder such as osteoarthritis or an undiagnosed posterior dislocation of the shoulder.

**Instability**

*Anterior dislocation* is a common sports injury. The anterior capsule and labrum are torn off the glenoid rim, and the humeral head is held anteromedially deep to the glenoid. It may reduce spontaneously, in which case the diagnosis can be difficult, or remain dislocated due to the pain and consequent muscle spasm. The diagnosis is then obvious, for not only is the shoulder held immobile, but the absence of the head from the socket also unmask the lateral prominence of the acromion. If, following reduction, the capsular tear does not heal, recurrent subluxation/dislocation is likely to occur. *Posterior dislocation* is fortunately rare, since it is difficult to diagnose as there is no gross deformity since the humeral head moves posteriorly to lie directly under the acromion. Close inspection from above the shoulder is the best way to observe the altered outline. The smooth, rounded prominence of the anterior aspect of the head is missing, and instead there is a fullness posteriorly. The deception is increased by the following: (i) there may be no history of injury, since posterior dislocation can occur in an epileptic fit – these patients may not present for a few weeks. As the hallmark is a gross restriction of movement, especially internal rotation and abduction, so an erroneous diagnosis of frozen shoulder can easily be made; and (ii) anteroposterior X-rays appear to show the head in correct alignment with the glenoid (Fig. U.9). An axillary view will confirm that the head is lying posteriorly.

The integrity of the axillary nerve should be established prior to reducing any dislocation. Failure to do so may result in medicolegal litigation. Since the deltoid will be inhibited by the pain of the dislocation, carefully examine the sensory distribution of the nerve, which is in a small area on the lateral aspect of the shoulder over the distal part of the deltoid.

In many instances, the dislocation, whether anterior or posterior, reduces spontaneously, but the patient may be left with instability and recurrent weakness in performing certain movements. While this diagnosis is easy to suspect, it can be difficult to prove. Furthermore, instability must be differentiated from

![Figure U.9](image-url) Posterior dislocation of the shoulder. (a) Anteroposterior X-ray showing ‘false congruity’. In fact, the humerus is lying directly posterior to the glenoid. The proximal humerus shows the ‘light-bulb appearance’ due to the altered rotation. (b) This posterior dislocation occurred during an epileptic fit and was initially treated as a frozen shoulder. The axillary view confirms that there is an impaction fracture of the humeral head on the posterior rim of the glenoid.
normal laxity. It is therefore essential to examine and compare both shoulders. The classical test for anterior dislocation is the apprehension test (Fig. U.10). The arm is placed at 90° of abduction and maximum external rotation, with the elbow flexed to 90°. The examiner then stabilizes the scapula with one hand, while increasing the external rotation abduction force with the other hand. If the shoulder is unstable, the patient will resist the movement, and also recognize that this is the position in which the shoulder ‘pops out’. To distinguish between instability and impingement, the shoulder is placed in a similar position to the apprehension test (i.e. abduction and external rotation) but, this time, with the patient lying supine, place one hand over the head of the humerus to prevent its anterior displacement. With instability, there is now no apprehension and no muscle contracture on further externally rotating with the examiner’s other hand; however, if there is impingement, the pain will still be reproduced. Multidirectional instability is due to a combination of anterior inferior plus posterior instability and is often associated with general ligamentous laxity. The sulcus sign is a useful test. This is done by distracting the humerus out of the glenoid by applying longitudinal traction – that is, with the patient upright or supine, pull on the arm. As the humeral head moves inferiorly, so a dent or a sulcus appears below the acromion. A gap of more than 1 cm is abnormal.

**Arthritis**

Rheumatoid arthritis of the glenohumeral joint is variable both in its severity and in the pattern of presentation. Three clinical pictures are recognized: the ‘dry form’, which presents as stiffness and bony crepitus secondary to loss of the joint space; the ‘wet form’, with marked synovial reaction; and the ‘resorptive form’, in which there is marked bone erosion of both the glenoid and the humeral head. Primary osteoarthritis was, until recently, uncommon in the shoulder, but it is now increasing in frequency due to the longevity of the population – most cases present over the age of 75 (Fig. U.11). Osteoarthritis below this age is nearly always secondary to previous injury (e.g. dislocation and fractures involving the humeral head) or to massive unreparable tears of the rotator cuff, which allow the humeral head to migrate proximally and eventually articulate against the underside of the acromion. The shoulder is the second most common site after the hip to be affected by osteonecrosis (Fig. U.12). This reflects the fragile vascular supply to the large

**Figure U.10** (a) Full external rotation in abduction is present in the right shoulder. (b) In the unstable left shoulder, this movement is resisted – the ‘apprehension’ test for anterior subluxation. The patient’s facial expression changes from a confident smile with the right arm to an apprehensive grimace on the left.

**Figure U.11** Osteoarthritis of the shoulder in a man aged 85 years.

**Figure U.12** Marked subchondral sclerosis of the humeral head secondary to avascular change in sickle-cell disease.
humeral head. In contrast to the hip, most cases have a well-defined precipitating cause. High-dose steroid treatment, for example as used in the treatment of head injuries, appears to have a particular affinity for precipitating osteonecrosis in the shoulder. It is also the most commonly affected joint in Caisson disease (the ‘bends’ in deep-sea divers and tunnel workers). MRI is the investigation of choice, especially in the early stages when plain X-rays are normal.

**Infection**

Both the glenohumeral joint and the humerus are uncommon sites for *acute pyogenic infection*. As a consequence, there may be a delay in making the diagnosis, which is often falsely attributed to trauma – in particular, fracture separation of the proximal humeral epiphysis due to a misinterpretation of the normal X-ray. (An antero-posterior film taken in slight rotation produces an appearance of a slight offset step between the metaphysis and the epiphysis at the level of the epiphyseal plate.) *Tuberculous infection* of the shoulder is also uncommon. It is easily mistaken for a frozen shoulder, from which it remains an important but little-known differential diagnosis. The historic name ‘caries sicca’ (dry tuberculosis) is a testament to diagnostic confusion in an earlier generation, with most cases probably being a frozen shoulder rather than tuberculosis!

**Tumours and tumour-like conditions**

These are discussed under LOWER LIMB, PAIN IN ‘Pain in and around the knee’ and ‘Tumours of bone’ (pp. 362 and 371, respectively).

**Miscellaneous**

*Rupture of the tendon of the long head of the biceps* affects elderly men and usually occurs after lifting (Fig. U.13). Something is felt to snap in the shoulder. There is usually some bruising in the upper arm, but the most dramatic sign is that the belly of the muscle rolls up into a ball when the biceps is tensed. Patients often think they have developed a tumour. Occasionally, the biceps can avulse from its distal attachment to the radial neck. *Bicipital tendinitis* presents in two ways: either in association with a rotator cuff impingement; or as an isolated condition in young adults. In the latter, tenderness is localized to the bicipital groove and aggravated by resisted active use of the biceps and resisted supination of the forearm with the elbow flexed to 90°.

**PAIN IN THE ELBOW**

Pain in the elbow is often well localized and characteristic, for instance a tennis elbow. However, there is frequently dual pathology – for example, cervical spondylosis with referred pain plus stiffness in the wrist causing abnormal use and stresses on the elbow, etc. A full history and examination of the upper limb plus the cervical spine including the peripheral neurology should be routine. On examination, check for any deformity (in particular the carrying angle) by comparing both elbows fully extended, or in comparable positions if extension is limited. Swellings of the olecranon bursa and subcutaneous nodules from rheumatoid arthritis or gout are usually obvious, but synovial thickening and an effusion can be easily overlooked. This is best checked by inspecting and feeling the soft tissues on each side of the olecranon. The joint line can be easily found on the lateral side. Feel just distal to the lateral epicondyle while pronating and supinating the forearm to ‘outline’ the radial head. On the medial side, the joint space is obscured by the muscle mass of the forearm flexors arising from the medial epicondyle. The ulnar nerve is easily palpated behind the medial epicondyle. Check the alignment of the three bony points – with the elbow flexed to 90°, the olecranon lies distally at the apex of a triangle formed with the two epicondyles; compare flexion and extension in both elbows. Since the elbow joint works in mid-range for most activities, many patients do not actually notice that they have lost 20–30° of extension. Full extension is useful for carrying heavy objects such as a bucket, but only seems essential for bowling in cricket! Check pronation and supination, which is often markedly reduced by relatively minor incongruity between the head of the radius and the capitulum. *Epicondylitis* is considered first as it is an extremely common symptom (Box U.5). The name itself is a misnomer, as not only is there no inflammation but also the problem is situated slightly more distally in origins of the extensor or flexor muscles.

![Figure U.13 Rupture of the right biceps (arrowed). The outline of the biceps is exaggerated, and there is a deep sulcus proximally.](image)
Tennis elbow affects 1–3 per cent of the total population in the 40- to 50-year age bracket and, if one considers all ages, the incidence rises to 20 per cent of men and 10 per cent of women. At least 50 per cent of patients ‘put up’ with their symptoms and do not trouble their doctor. It is caused by overuse of the extensors, particularly extensor carpi radialis brevis – classically the backhand stroke at tennis. The pain comes on gradually, often after a period of unaccustomed excessive exercise. It usually localizes to the lateral epicondyle but, when severe, it can radiate widely. It is aggravated by simple activities that use the forearm in pronation, for example pouring tea from a teapot. As with other tendonitis, there is a triad of signs: tenderness just below the lateral epicondyle; pain on resisted active extension of the wrist; and pain on passive stretch (i.e. combined pronation and palmar flexion of the wrist with the elbow at 90°). X-rays are usually normal, but occasionally calcification is seen at the extensor origin. Rarely, this calcification may precipitate an acute calcification syndrome, which is extremely painful and is accompanied by marked inflammation with warmth and swelling over the lateral side of the elbow, to the extent that it can be easily mistaken for an acute severe infection. Other causes of lateral elbow pain are entrapment of the posterior interosseous nerve between the two heads of the supinator muscle at the level of the radial neck (see ‘Nerve entrapments’, below), and osteoarthritis affecting the radiocapitular joint. With the latter, pronation/supination is restricted. Golfer’s elbow is similar to tennis elbow but involves the flexor origin from the medial epicondyle. The diagnosis is often less clear-cut, and it can be difficult to distinguish from an ulnar neuritis or sprains of the medial ligament. The triad of signs are: local tenderness just distal to the epicondyle; pain on resisted palmar flexion of the wrist; and pain on combined supination and extension of the wrist.

**Arthritis**

Rheumatoid arthritis commonly affects the elbow; approximately 75 per cent of patients complain of some pain, and the joint is severely involved in 20 per cent. X-ray features vary from osteoporosis and soft-tissue swelling in the early stages, to gross bone loss of the distal humerus with erosion into the olecranon in later stages, when a fracture secondary to a fall can easily result in a flail elbow. The ulnar nerve is at risk both from the early synovitis and the deformity secondary to the destructive arthropathy. The surprising feature about osteoarthritis is that, while it frequently involves the elbow, many patients are unaware that they have a problem since they are not inconvenienced by loss of 20–30° of flexion provided they can get their hand up to eat, wash and comb their hair. Loss of supination is more troublesome than an equivalent loss of pronation as the latter can be compensated for by abduction of the shoulder. The elbow is the second most common joint (after the knee) to be affected by haemophilic arthropathy; this may eventually lead to severe ankylosis. The most common cause of a very stiff and painful elbow is traumatic osteoarthritis secondary to severe intra-articular fractures.

**Infection**

Symptoms and signs of an acute septic arthritis are similar to those described for the knee – fever, synovial thickening, warmth, tenderness and immobility. The X-rays are normal, and the diagnosis is made on aspirating the joint (see LOWER LIMB, PAIN IN ‘Pain in and around the knee’, p. 362). Osteomyelitis can affect the metaphyseal area of any of the three bones, but it is much less common than in the lower limb. There is frequently a delay in diagnosing both tuberculous osteitis and synovitis. Furthermore, a discharging sinus is easily mistaken for a discharging olecranon bursa if it points at the back of the elbow.
Loose bodies/locked elbow
The elbow is the second most common joint, after the knee, for loose bodies. The convex surface of the capitulum is one of the classic sites for osteochondritis dissecans. This can present with lateral elbow pain or, if the fragments separate as a loose body, can cause locking. It occurs in adolescents, but it may remain asymptomatic until later in life when the elbow locks. An X-ray usually reveals two or three significantly sized loose bodies. The reason for the time delay is that the loose bodies, by deriving their nutrition from the synovial fluid, can continue to grow in size, eventually becoming large enough to limit movement. Multiple loose bodies are due to synovial chondromatosis (Fig. U.14), which is now regarded as a benign synovial neoplasm (see LOWER LIMB, PAIN IN ‘Pain in and around the knee’, p. 362).

Instability
The elbow is particularly prone to trauma; it is the second most common joint, after the shoulder, to suffer dislocation. Dislocations, fracture dislocations (especially fractures of the coronoid or the radial head) and fractures (the radial head in combination with the medial ligament injury) can result in major instability. Elbow injuries are deceptive; in particular, dislocation can spontaneously reduce, and small bony avulsions are easily missed on X-rays. Always be suspicious of a bruised swollen elbow, and check for any neurovascular complications. A good example is entrapment of the medial epicondyle together with the ulnar nerve within the joint in a child (see ‘Nerve entrapments’, below). The main causes of neuropathic arthropathy are tabes dorsalis, syringomyelia, diabetes and congenital indifference to pain. Tabes dorsalis predominantly affects the lower limbs, while syringomyelia involves the upper limbs. Diabetes particularly affects the foot. Charcot joints can be painful in their early stages before the joint disintegrates. Excessive osteophyte formation in the presence of gross instability is the hallmark of a Charcot joint.

Sprains of the medial collateral ligament are an occupational hazard of baseball players, in particular pitchers.

Nerve entrapments
The ulnar nerve can be compromised at several anatomical sites around the elbow: (i) in the intermuscular septum as the nerve passes from the anterior to the posterior compartment of the arm; (ii) in the cubital tunnel behind the medial epicondyle, where it is vulnerable from direct injury, or pressure from synovitis of the elbow in rheumatoid or osteophytes in osteoarthritis; and (iii) where it passes between the two heads of flexor carpi ulnaris, just distal to the elbow. In addition, there are two specific childhood fractures. The childhood lateral condylar fracture of the distal humerus, if poorly treated, can leave a valgus deformity (increase in the carrying angle), which may eventually cause an ulnar nerve paresis many years later (Fig. U.15). The other injury causes acute ulnar nerve symptoms. In a dislocation of the elbow with an avulsion of the medial epicondyle, the joint often spontaneously relocates, but the medial epicondyle – together with the flexor origin and the ulnar nerve – may become trapped within the elbow joint. Since the small bony fragment is obscured by the overlying bone, this injury and its complication can be easily missed by the unwary. A comparative X-ray of the opposite elbow will confirm that the epicondyle is missing.

Figure U.14 Synovial chondromatosis of the elbow with multiple loose bodies (arrowed).
In the early stages of ulnar neuritis, symptoms predominate over signs. The patient presents with a history of paraesthesia and numbness that, on close questioning, affect the ring and little fingers. The first sign is a loss of sweating (e.g. dryness of the little finger and ulnar half of the ring finger). Continuing compression or irritation leads to weakness in using the hand. Look for wasting of the hypothenar eminence, weakness of abduction and adduction of the fingers and loss of power of pinch. Muscle wasting is best seen and felt in the web space between the thumb and index metacarpals (adductor pollicis and the first dorsal interosseous muscles, both of which are supplied by the ulnar nerve). Due to the muscle loss, the pinch grip is affected and the thumb collapses into hyperextension at the metacarpophalangeal joint, with flexion at the interphalangeal joint (Froment’s sign, Fig. U.16).

The median nerve is particularly at risk with major elbow injuries, for example the supracondylar fracture of the humerus in children or a posterior dislocation of the elbow, since the nerve has a relative fixed point where it passes between the two heads of pronator teres. A chronic entrapment can also occur at this site. In addition to the altered sensation in the median nerve distribution in the hand, look for local tenderness by compressing the nerve as it passes under pronator teres while simultaneously resisting the patient pronating the forearm. Entrapment of the posterior interosseous branch of the radial nerve where it passes through the supinator muscle is difficult to distinguish from tennis elbow. The tenderness is situated more anteriorly over the radial neck and becomes localized to this site during resisted supination of the forearm. The patient may complain of weakness of grip due to the failure of the extensor muscles to stabilize the wrist. Electromyography (EMG) is useful in confirming the clinical diagnosis of a nerve entrapment, especially if there is symptomatic cervical spondylosis with nerve root irritation – the ‘double-hit’ phenomenon in which a nerve is irritated at two sites.

Swellings
Anatomically there are a considerable number of bursae around the elbow joint. The majority are deep, for example, the bursa between the biceps tendon and the radial tuberosity. Only the olecranon bursa is of clinical relevance, the common cause of bursitis being recurrent minor trauma. Often referred to as ‘student’s elbow’, it is most frequently seen in office workers – leaning on the elbow while using a telephone is a common culprit. It presents as a fluctuant, non-tender, clearly defined swelling without tenderness or other signs of inflammation and with full elbow mobility (Fig. U.17). If there are any signs of inflammation, look for an underlying diagnosis. The common inflammatory causes are rheumatoid arthritis, gout or infection; typically, 20–30 per cent of all olecranon bursitis will be infected. If there is any suspicion of infection, aspirate and culture the fluid. In the presence of chronic infection – particularly if there is a discharging sinus – an X-ray is required to check that there is no pathology such as tuberculosis within the joint or the bones.

Figure U.16 Froment's sign; testing for weak hand intrinsic muscles, in particular the 1st dorsal interosseous and adductor pollicis, as found in ulnar nerve palsy. Weakness of these means pinch grip is performed by the flexor pollicis longus with flexion of the interphalangeal joint of the thumb as seen on the right hand.

Figure U.17 A large olecranon bursa.
The posterior subcutaneous border of the ulna is a common site for inflammatory subcutaneous nodules. Rheumatoid arthritis and gout are the common cause.

**PAIN IN THE FOREARM**

Tendinitis and tenosynovitis are considered in the next section. The forearm is the second most common anatomical site, after the lower leg, for a compartment syndrome. In both situations, there are two bones, a tough interosseous membrane and well-defined compartments bounded by a strong overlying deep fascia. If undiagnosed, it leads to Volkmann's ischaemic contracture (Fig. U.18), in which loss of the forearm flexors to the fingers and thumb, together with loss of median and sometimes the radial nerve, results in a useless hand. Muscles supplying the wrist usually survive, but the fingers show a fixed length phenomenon – that is, palmar flexion of the wrist automatically causes extension of the finger at the interphalangeal and metacarpal phalangeal joints, while on dorsiflexion the fingers become clawed. Classically, the condition is due to a displaced supracondylar fracture of the distal humerus in a child, the brachial artery being occluded and damaged against the sharp distal end of the proximal fragment (i.e. across the metaphysis of the humerus). Other causes are crush injuries to the forearm; postoperatively after both vascular and orthopaedic surgery (there is a particular risk following plating of the radius); and bleeding disorders. The signs and symptoms are similar to those described for the lower limb. The cardinal symptom is pain, and the cardinal sign is pain on passive extension of the fingers. Other features include paraesthesiae progressing to numbness and paralysis, and a solid woody feel to the forearm on palpation. The pulse can be highly misleading as it may be transmitted down an artery that is not passing through the involved compartment.

**THE WRIST: PAIN AND DEFORMITY**

Causes of pain and deformity in the wrist are shown in Box U.6.

**Box U.6 Diagnostic categories of wrist pain and deformity**

<table>
<thead>
<tr>
<th>Deformity</th>
<th>Arthritis</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Congenital</strong></td>
<td>• Rheumatoid arthritis</td>
</tr>
<tr>
<td>• Radial club hand</td>
<td>• Osteoarthritis</td>
</tr>
<tr>
<td>• Ulnar club hand</td>
<td>– Wrist joint (usually secondary to trauma, e.g. scaphoid fracture; dislocation/ligament injuries causing carpal instability)</td>
</tr>
<tr>
<td>• Distal ulnar dysplasia</td>
<td>– Distal radio-ulnar joint</td>
</tr>
<tr>
<td><strong>Acquired</strong></td>
<td>• Osteonecrosis of lunate (Kienböck’s disease)</td>
</tr>
<tr>
<td>• Post-traumatic</td>
<td><strong>Infection</strong></td>
</tr>
<tr>
<td>– Growth plate injuries</td>
<td>• Acute – pyogenic</td>
</tr>
<tr>
<td>– Malunion (e.g. Colles’ fracture)</td>
<td>• Chronic – tuberculous</td>
</tr>
<tr>
<td>• Rheumatoid arthritis</td>
<td><strong>Nerve entrapments</strong></td>
</tr>
<tr>
<td>• Nerve injuries (e.g. wrist-drop)</td>
<td>• Carpal tunnel syndrome</td>
</tr>
<tr>
<td><strong>Carpal tunnel syndrome</strong></td>
<td>• Ulnar nerve – distal entrapment</td>
</tr>
<tr>
<td><strong>Conditions of tendons</strong></td>
<td><strong>Conditions of tendons</strong></td>
</tr>
<tr>
<td>• Tenosynovitis</td>
<td>• Tendinopathy</td>
</tr>
<tr>
<td>– Infective</td>
<td>– Inflammatory</td>
</tr>
<tr>
<td>– Inflammatory</td>
<td>– Mechanical</td>
</tr>
<tr>
<td>– Mechanical</td>
<td>(e.g. De Quervain’s)</td>
</tr>
<tr>
<td>(e.g. De Quervain’s)</td>
<td>• Tendon ruptures</td>
</tr>
<tr>
<td><strong>Miscellaneous</strong></td>
<td><strong>Miscellaneous</strong></td>
</tr>
<tr>
<td>• Ganglion</td>
<td>• Chronic wrist pain/carpal instability</td>
</tr>
</tbody>
</table>

**Figure U.18 Complications of a supracondylar fracture of the distal humerus.** (a) Volkmann’s ischaemic contracture of the forearm and the hand. Surgical decompression of the forearm muscles was carried out too late to prevent ischaemic necrosis. In addition, there was loss of both the median and ulnar nerves. (b) Varus deformity of the right elbow (the gun-stock deformity) due to a malunion.
Deformity
Congenital deformities are rare and complex. In general, they either are due to failure of formation of part of the skeleton or occur as part of a generalized disease. Radial and ulnar club hands are examples of ‘arrest of development’ in which a ray may be partially or completely absent (see Fig. S.49). For example, in a radial club hand, the thumb, first metacarpal, scaphoid and trapezium will be absent, and the radius will be only partially developed. Without an adequate wrist joint, the hand falls off the ulna and collapses into radial deviation. Failure of development on the ulnar side produces the opposite deformity. Distal ulnar dysplasia is an example of a generalized disease and is associated with multiple exostoses. Secondary to the ulnar defect, the radius is bowed, and the radial head is dislocated at the elbow.

Acquired deformities around the wrist are common. Displaced Colles’ fractures invariably heal with some deformity despite an initial satisfactory reduction. Premature fusion of the distal radial epiphysis occasionally occurs following a fracture separation in childhood. This can produce a marked deformity with a radial tilt and relative overgrowth of the ulna. Severe destruction of the wrist and carpal joints is a hallmark of rheumatoid arthritis. A ‘wrist-drop’ is of course not a true deformity but results from an injury to the radial nerve and paralysis of the extensor muscles – in order to grip, it is essential to be able to extend the wrist.

Nerve entrapments/injuries
The carpal tunnel syndrome, in which median nerve function is impaired beneath the carpal ligament, is the most common of all entrapment syndromes. Although in most cases it is idiopathic, it should not be considered a primary diagnosis since it may be secondary to systemic disease, such as diabetes and thyroid dysfunction, or local disease such as rheumatoid arthritis, which causes mechanical impingement into the carpal tunnel. It commonly occurs in pregnancy, but the symptoms are usually mild and transient, disappearing once the baby is born. The features are paraesthesia and numbness, most noticeable at night, affecting the median nerve distribution. The symptoms are relieved by shaking the hand or holding it over the side of the bed (Fig. U.19a). Prolonged and severe compression will eventually cause motor weakness – weakness of thumb opposition (Fig. U.19b). Frequently, no abnormal physical signs can be detected in the clinic. Occasionally, there is a positive Tinel’s sign over the carpal tunnel, and sometimes paraesthesiae can be precipitated by keeping the wrist in full palmar flexion for a minute.

Wasting of the thenar eminence is a late feature. An EMG is particularly useful, either when the diagnosis is not clear-cut or when there are associated radicular problems in the arm secondary to cervical spondylosis. Compression of the ulnar nerve in the palm of the hand in the carpal tunnel is a well-described (although uncommon) condition. Diagnostically, it can be confusing since, depending on the precise location of the compression in relation to the division of the nerve into a superficial sensory branch and a deep motor branch, the signs may be purely sensory, purely motor or a mixture of both. Isolated motor loss with a gradual onset of clumsiness in the hand is easy to miss. Ulnar clawing of the ring and little fingers is much more obvious in distal ulnar nerve injuries than in proximal injuries around the elbow, as the clawing will be less marked when the medial ulnar half of the flexor digitorum profundus and the distal small muscles to the ring and little finger are both paralysed.

Tendons
Acute pyogenic infection of the flexor tendon sheaths is considered later (see ‘Conditions affecting the hands and fingers’). An important anatomical point is that the synovial sheath of flexor pollicis longus extends from the tendon insertion on the distal phalanx proximally through the carpal tunnel into the distal forearm. The common sheath of the digital flexors also passes under the carpal ligament into the palm, and then it continues down into the flexor sheath of the little finger. The extensor tendon sheaths extend distal and proximal to the extensor retinaculum but do not extend down into the fingers. Acute infection of the
Extensor sheath inflammation is usually due to a penetrating injury. There is often an associated overlying cellulitis, which can obscure infection in the underlying sheath.

Inflammatory tenosynovitis is common in rheumatoid disease and affects both the flexor and the extensor tendons (Fig. U.20). The thickened swollen synovium is more obvious around the dorsal extensor tendons than the palmar flexor tendons, where the swelling is confined to the proximal palm and the distal forearm by the constriction of the tight carpal ligament. Pain on flexion of the fingers in the presence of good individual finger function is the main feature of the rheumatoid flexor tenosynovitis at the wrist. Extensor tendon ruptures in rheumatoid are not only common but also easily overlooked by both the patient and the doctor in a severely affected rheumatoid hand. They present with a ‘dropped finger’; this is an inability actively to extend the metacarpophalangeal joint. Due to the action of the interossei muscles, extension at the proximal interphalangeal joint is retained, hence the deception. Rupture is due to two factors: rheumatoid synovium from the tendon sheath invading into the tendon; and abrasion over the ulnar head. Failure to make the diagnosis early will lead to sequential dropping of the fingers and an unnecessary additional loss of function in the hand.

Some ruptures are mechanical, for example, rupture of the extensor pollicis longus approximately 6 weeks after a minimally displaced Colles’ fracture, which results in loss of extension of the distal phalanx of the thumb. Mechanical or stress-induced tenosynovitis around the wrist is common, usually following a period of repetitive unaccustomed activity, for example an office worker using a pick-axe and causing De Quervain’s syndrome (tenosynovitis of the tendon sheath of extensor pollicis brevis and abductor pollicis longus where they pass through a tunnel over the radial styloid). The triad of signs is local tenderness over the radial styloid, pain on resisted ulnar deviation with the thumb abducted, and pain on passive stretch into radial deviation with the thumb first placed fully opposed across the palm. Differential diagnosis is from a fractured scaphoid and carpometacarpal osteoarthritis between the trapezium and the thumb metacarpal.

**Arthritis**

The wrist joints are the second most commonly involved joints in rheumatoid arthritis, after the metacarpophalangeal joints (see Fig. U.20). Initial involvement is often on the ulnar side, especially the distal radio-ulnar joint. As the disease progresses into the radial carpal and intercarpal joints, so the wrist develops the classical deformity with radial deviation and palmar subluxation. As mentioned above, the synovial sheaths of the tendons are frequently involved. Without a well-aligned balanced wrist, it is impossible to have a good finger grip. In contradistinction to rheumatoid, osteoarthritis of the wrist is uncommon unless there has been a previous intra-articular fracture or dislocation. Osteonecrosis affects the proximal convex surface of the lunate. There is usually an associated relative shortening of the ulna. The condition presents in early adult life with discomfort following exercise. Grip strength is usually diminished. In the early stages, the X-rays are normal, followed in time by sclerosis, then collapse of the articular surface and finally osteoarthritic change in the wrist joint (Fig. U.21). The proximal pole of the scaphoid is at risk following fractures across the proximal part of the wrist (Fig. U.22). The long-term result is osteoarthritis of the wrist.

**Figure U.20** (a) Hands in rheumatoid arthritis. (b) Note the swan-neck deformities of the fingers on the lateral views.
Infection

Acute pyogenic infection of the wrist joint is uncommon unless there has been a penetrating injury. However, the wrist is a common site for tuberculous infection, which often presents relatively late as there is a gradual onset of pain and stiffness. With progressive destruction, there is palmar subluxation, and it is easy to mistake for monoarticular rheumatoid. There is usually marked wasting of the forearm muscles. Involvement of the flexor sheath leads to large fluctuant swellings in the palm and distal anterior forearm, the two being connected under, but constricted by, the carpal ligament (a compound palmar ganglion). In the early stage, tuberculous arthritis shows local osteoporosis with pencilling of the cortical margins. With progression, there is bone erosion, leading eventually to destruction of the carpus and the wrist.

Tumours and tumour-like conditions

For a discussion on benign tumours, cysts of bone, tumour-like conditions and primary malignant bone tumours, see LOWER LIMB, PAIN IN (p. 371). The distal radius is the third most common site for an osteoclastoma (giant-cell tumour). This presents with aching discomfort and bony swelling of the distal radius. On plain X-rays, these tumours have a characteristic appearance as the cystic lesion extends up to the joint margin. It affects patients in the 20- to 40-year age group.

Miscellaneous

The most common site for a ganglion is the dorsum of the wrist, where they arise as a result of mucoid degeneration of the joint capsule. The patient is usually a young adult but may be middle-aged. Ganglia are rarely (if ever) seen in the elderly – that is, they eventually resolve. Many simply present with a painless lump. Symptoms are usually caused by pressure on adjacent structures; for example, a ganglion on the palmar aspect of the wrist may be associated with a carpal tunnel syndrome or ulnar nerve compression. On examination, there is a non-tender, well-defined smooth lump that is usually slightly fluctuant but may appear hard if it is small and beneath the deep fascia.

Chronic wrist pain is a complex subject. In brief, recent advances in MRI imaging and arthroscopy have revealed that the carpus is the site of complex
ligamentous injuries and tears. The triangular fibrocartilage, which binds the radius and ulna together but separates the inferior radio-ulnar joint from the wrist joint, can be torn rather like a meniscus in the knee. The most common type of carpal instability is \textit{scapholunate disassociation} secondary to disruption of the ligaments connecting the scaphoid to the lunate. This injury can occur following relatively minor ‘sprains’ of the wrists. Plain X-rays show widening of the gap between the scaphoid and the lunate (Fig. U.23a) on the anteroposterior film, and collapse of the normal alignment on the lateral film – the longitudinal axis of the radius, the capitate and third metacarpal should form a straight line. Patients present with pain, or weakness of the wrist and diminished grip, while certain movements may cause a click or a snap. These injuries are the cause of osteoarthritis later in life (Fig. U.23b).

**CONDITIONS AFFECTING THE HANDS AND FINGERS**

The hands and the fingers are affected by many systemic diseases (see \textit{JOINTS, AFFECTIONS OF}, p. 319, which deals with the generality of joint diseases). This section is concerned with conditions local to the hands or fingers and the local manifestation of generalized disease (Box U.7).

**Deformity**

This is a complex group of conditions that may be local to the hand or may occur as part of a generalized skeletal abnormality. \textit{Syndactyly} is the most common congenital hand deformity. The fingers and thumb may be joined by complete or partial webs of skin.

![Figure U.23](a) Scapholunate disassociation. Note the gap between the scaphoid and the lunate. (b) Osteoarthritis of the wrist is the long term complication.

<table>
<thead>
<tr>
<th>Box U.7</th>
<th>Diagnostic categories affecting the hand and fingers</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Deformity</strong></td>
<td><strong>Septic arthritis</strong></td>
</tr>
<tr>
<td><em>Congenital</em></td>
<td><em>Osteoarthritis</em></td>
</tr>
<tr>
<td>• Lesions localised to the hand</td>
<td><em>Necrotizing fasciitis</em></td>
</tr>
<tr>
<td>- Syndactyly</td>
<td><em>Deep palmar space infections</em></td>
</tr>
<tr>
<td>- Duplication</td>
<td><strong>Acquired</strong></td>
</tr>
<tr>
<td>- Macro/Microdactyly</td>
<td><em>Tuberculous</em></td>
</tr>
<tr>
<td>• Lesions associated with</td>
<td><em>Leprosy</em></td>
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<tr>
<td>generalised diseases</td>
<td><em>Fungal</em></td>
</tr>
<tr>
<td>(e.g. skeletal dysplasias)</td>
<td><em>Infection from bites</em></td>
</tr>
<tr>
<td><strong>Conditions of tendons</strong></td>
<td><strong>Arthritis</strong></td>
</tr>
<tr>
<td><em>Locking</em></td>
<td><em>Inflammatory</em></td>
</tr>
<tr>
<td>• Trigger finger</td>
<td><em>Rheumatoid arthritis</em></td>
</tr>
<tr>
<td>• Trigger thumb: children</td>
<td><em>Gout</em></td>
</tr>
<tr>
<td>(congenital), adults</td>
<td><em>Psoriatic arthropathy</em></td>
</tr>
<tr>
<td><strong>Ruptures</strong></td>
<td><strong>Osteoarthritis</strong></td>
</tr>
<tr>
<td>• Mallet finger</td>
<td><em>Heberden’s nodes</em></td>
</tr>
<tr>
<td>• ‘Dropped’ finger</td>
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- **Figure U.23** (a) Scapholunate disassociation. Note the gap between the scaphoid and the lunate. (b) Osteoarthritis of the wrist is the long term complication.
Dupuytren’s contracture affects any digit. There are a number of different types that have different incidences in different populations. For example, in the black American population, replication of the little finger is very common. Macrodactyly can affect any of the fingers or the thumbs and is secondary to a local hamartomatous enlargement. Microdactyly is the reverse.

Acquired deformities are common, most being due to local trauma, for example failure to regain movement in a digit following a laceration of a tendon, rotational malalignment of the finger secondary to a fracture, etc. An important group is contracture secondary to neurological disease, for example cerebral palsy, or an ulnar clawed hand secondary to a laceration of the ulnar nerve (see the earlier discussion on high and low ulnar nerve injuries). Dupuytren’s disease, which is due to contracture of the palmar fascia (Fig. U.24), is common in Northern Europe and rare in Asia. It is probably an autosomal dominant condition, but other risk factors are involved, such as smoking, alcohol, diabetes and (possibly) epilepsy. The ring and little fingers are the most commonly affected. In the early stages, there is a thickening in the palm, which manifests itself as firm nodules with puckering and pitting of the overlying skin. With time, this develops into inextensible thickened bands, which usually run up into the fingers, causing loss of extension of the metacarpophalangeal joint and contractures of the interphalangeal joints. In severe cases of Dupuytren’s disease, the fingernails will be held impacted down into the palm. Knuckle pads may be present over the dorsum of the proximal interphalangeal joints.

The condition can also affect the feet (the plantar fascia) and may occasionally be associated with fibrosis of the corpus cavernosum (Peyronie’s disease). In early cases, the differential diagnosis is from an implantation dermoid cyst, which causes a more superficial nodule in the skin and a contracture of a flexor tendon in which, contrary to Dupuytren’s, the tight band moves on passively flexing the finger.

Tendons
‘Triggering’ can affect any finger including the thumb, but it is most frequently seen in the middle and ring fingers. Flexion (often accompanied by a click) is full but, on trying to extend, the finger remains stuck until the patient applies an extra active extension effort, or unlocks the finger passively with the other hand. Most cases arise spontaneously due to thickening in the flexor tendon, accompanied by thickening and stenosis at the entrance to the fibrous flexor profundus tunnel in the palm. However, it is important to exclude associated underlying disease such as diabetes, rheumatoid arthritis, gout or amyloid. Congenital trigger thumb is a well-recognized entity in children, although there is dispute as to whether it is really congenital, as it is rarely (if ever) seen before the age of 6 months.

Mallet finger results from a loss of active extension at the distal interphalangeal joint secondary to a traumatic avulsion of the extensor tendon at its insertion into the base of the distal phalanx. It is a common injury in cricketers and baseball players, where the hard ball can catch the tip of the finger, forcing it into sudden flexion. While passive extension is present, there is a loss of active extension. A lateral X-ray will distinguish between a pure tendon rupture and an avulsion fracture. The most common cause of a dropped terminal phalanx of the thumb is a traumatic rupture of extensor pollicis longus at the wrist (see ‘The wrist: pain and deformity’, p. 713).

Boutonnière (buttonhole) deformity is due to a rupture or attenuation of the central slip of the extensor hood at its insertion onto the base of the proximal phalanx. As the name implies, the proximal interphalangeal joint buttonholes through the resulting defect, the adjacent extensor slips being pushed aside. This allows an excessive extensor pull at the distal interphalangeal joint so that, in the final deformity, the finger is fixed flexed at the proximal interphalangeal joint and in hyperextension at the distal joint. The reverse deformity to a boutonnière is the swan-neck. In which there is extensor overactivity at the proximal interphalangeal joint and a simultaneous flexion deformity at the distal joint. There are a number of different types of swan-neck deformity.
of causes, including contracture of the intrinsic muscles acting at the proximal interphalangeal joint, mallet finger deformity at the distal phalangeal joint, and paralysis or division of flexor digitorum superficialis, which causes weakening of the flexor action at the proximal interphalangeal joint. As with the boutonnière, the swan-neck is commonly seen in a rheumatoid hand (see Fig. U.20). Non-infective tenosynovitis occurs with inflammatory disorders such as rheumatoid arthritis, and occasionally from unaccustomed repetitive overuse of a finger. It is only seen with flexor tendons, since the extensor tendons do not have synovial sheaths in the fingers.

**Infections**

Infections in the hand and fingers are common, important but often poorly diagnosed and managed. It is all too easy to confuse the common superficial infections with the destructive, although fortunately less frequent, deep infections. Furthermore, due to the tough compact nature of the skin and subcutaneous tissues of the palm and palmar surface of the fingers, the swelling of inflammatory oedema appears on the back of the hand, thereby masking the anatomical site of origin of many deep infections. The serious complication of lymphangitis and septicæmia can rapidly supervene. Therefore, as well as assessing the hand, always take the temperature and feel the regional lymph nodes.

In examining the hand, not only look for signs of inflammation, but also think anatomically. For example, a deep infection involving the little finger will spread along the tendon sheath — that is, through the palm, under the carpal tunnel and up into the forearm. A full medical history and examination are required to exclude diseases such as diabetes, steroids, HIV and other immunocompromising conditions. Check the tetanus status. A full blood count should be performed, and cultures obtained prior to starting antibiotic treatment. The ESR and CRP can be useful in monitoring the response, especially to starting antibiotic treatment. The ESR and CRP can be useful in determining the presence of osteomyelitis or the spread of infection into the subfascial space. Due to the tough overlying palmar skin, signs of inflammation and swelling are most obvious on the dorsal aspect of the hand in the interval between the adjacent metacarpals. An abscess in this region may be of the collar stud variety — that is, the pus pointing on the dorsum, but the main abscess lying much more anteriorly, deep to the palmar skin. Deep palmar space infections are fortunately uncommon. They form in the space between the palmar aponeurosis and the third, fourth and fifth metacarpals, which is traversed by the flexor tendons to the little, ring and middle fingers, the appropriate digital nerves and vessels, and the superficial palmar arch. The hand is markedly swollen, and in particular the concavity of the palm is lost. If this sign is present, do not be deceived by the excessive swelling on the dorsum.

**Cellulitis** is an infection of the skin and subcutaneous tissues. It is very important to distinguish a true cellulitis from the local signs of infection in the skin overlying an abscess. Necrotizing fasciitis, in its early stages, often presents as a widespread, very painful cellulitis with marked systemic symptoms. Fortunately, it is rare, but it carries a high mortality. Drug addicts, immunocompromised patients and those with diabetes are most at risk from this form of severe cellulitis.

**Paronychia (a nailfold infection)** is the most common of all hand infections. When acute, there is erythema, swelling and tenderness around the nailfold and extending across the dorsum of the distal phalanx over the nail bed. Pus may be released by gently lifting the nail. Chronic paronychia requires a bacterial diagnosis as it may be due to the persistence of pyogenic infection, to a fungal infection, or occasionally to tuberculosis. An X-ray should be taken to rule out chronic osteomyelitis. A felon or whitlow is a finger pulp infection, and presents with a swollen, red and acutely tender fingertip. Infective tenosynovitis is usually secondary to a puncture wound and involves the fibrous flexor sheath of a finger. The ring, middle and index fingers are the most commonly affected; in these fingers, the infection is limited to the finger and palm distal to the distal palmar skin crease (i.e. distal to the metacarpophalangeal joint). The finger will be painful, swollen, held rigidly semi-flexed and very tender, and any movement will be firmly resisted. Infections involving the tendon sheaths of the thumb and little finger will extend through the palm under the carpal ligament and into the distal forearm.

**Web space infections** often start as abrasions, blisters or callus on the distal part of the palm, with infection spreading into the subfascial space. Due to the tough overlying palmar skin, signs of inflammation and swelling are most obvious on the dorsal aspect of the hand in the interval between the adjacent metacarpals. An abscess in this region may be of the collar stud variety — that is, the pus pointing on the dorsum, but the main abscess lying much more anteriorly, deep to the palmar skin. Deep palmar space infections are fortunately uncommon. They form in the space between the palmar aponeurosis and the third, fourth and fifth metacarpals, which is traversed by the flexor tendons to the little, ring and middle fingers, the appropriate digital nerves and vessels, and the superficial palmar arch. The hand is markedly swollen, and in particular the concavity of the palm is lost. If this sign is present, do not be deceived by the excessive swelling on the dorsum.

**Deep thenar space infection** occurs in the potential space between the palmar aponeurosis and the fascia overlying adductor pollicis, which contains not only the flexor pollicis longus tendon plus the neurovascular bundles to the thumb but also the neurovascular
bundle on the radial side of the index plus the flexor tendons of the index. Therefore, a deep thenar space infection can involve both the thumb and the index finger.

Acute pyogenic septic arthritis may result from a penetrating wound or be a complication in rheumatoid arthritis – an acute onset of inflammation in a single joint in a multiply involved rheumatoid hand should alert the physician to this complication. Differential diagnosis in an otherwise normal hand is from gout. Osteomyelitis is easy to miss in the hand as it is mistaken for an underlying soft-tissue infection. It can also occur as a result of poorly treated soft-tissue or joint infections. The usual error is to omit to take an X-ray.

Tuberculosis presents with a variety of pathologies – most commonly a chronic tenosynovitis – but it can cause osteomyelitis or a joint synovitis. A biopsy is usually required for diagnosis. Other mycobacteria may be encountered, for example Mycobacterium marinum from seawater. Leprosy (Mycobacterium leprae) usually presents with trophic changes in the hands and feet.

Fungal infections
There are a wide variety of fungal infections, which can involve any of the tissues or anatomical spaces discussed under pyogenic infections. Therefore, these must be included in any differential diagnosis for a subacute or chronic infection. Diabetic and immunocompromised patients are particularly at risk. Unfortunately, the diagnosis is often delayed as fungal infections tend to be overlooked. They are particularly common in chronic paronychia, chronic infection of the nailplate (onychomycosis) and granulomas, especially if the original wound was from a thorn. When fungi cause deep infections (fortunately uncommon), the organism can be difficult to isolate. Mycetomas (Madura ulcer) are due to subcutaneous implantation of fungal spores or actinomycosis. These can affect the fingers and present with pseudotumours and discharging sinuses (see Nodules, ‘Fungi’, p. 465).

Bites
One of the common wounds to a hand is a bite, the most frequent culprit being a dog. Fortunately, most wounds do not become infected but, when they do, a wide variety of organisms can be involved. Cat bites, while less frequent, have a greater tendency to infection. Cat scratch fever can be a further complication. Bites from animals are also an entry point for rabies. Human bites are unpleasant and have a particular propensity to infection, usually by several species of organisms. Most are caused by fighting between children; when they result from adult domestic violence, they can be deceptive, as the recipient often fails to give a full history. Venomous bites and stings from snakes, bees, spiders and other insects may, in addition to systemic effects, often cause widespread local or regional inflammation, which can progress to local necrosis and which in turn may become secondarily infected. As the onset is so acute, there is usually a clear history, but occasionally bee or mosquito stings can be confused with a cellulitis.

Arthritis
The hand is involved in many inflammatory arthropathies. The peculiar feature of rheumatoid arthritis is the sparing of the distal interphalangeal joints, whereas the other small joints of the hand and the carpus are almost invariably involved. The disease often presents with early morning stiffness affecting, in particular, the metacarpophalangeal joints and the proximal interphalangeal joints. In the early stages, there is a synovitis with swelling, warmth, tenderness, stiffness and painful movement of some or all of these joints. In long-standing disease, the classical deformities of ulnar drift at the metacarpophalangeal joints with progressive palmar subluxation and eventual dislocation of the fingers become evident. Swan-neck and boutonnière deformities are particularly destructive of finger function.

It is easy to overlook ‘dropped fingers’; the initial finger affected is usually the little finger. The cause is extensor tendon rupture in the region of the inferior radio-ulnar joint due to a combination of synovitis and bony impingement. If the diagnosis is missed, there can be a sequential ‘dropping’ of all the other fingers. Involvement of the thumb causes a loss of pinch grip as the thumb assumes a swan-neck or boutonnière type of deformity – there is either fixed flexion of the carpometacarpal joint with hyperextension at the interphalangeal joint, or vice versa. The extent of bone and joint destruction on X-rays is usually greater than one would expect from the clinical examination. Early features are soft-tissue swelling, periarticular erosions and osteoporosis, followed later by loss of joint space, bony destruction and joint disintegration, subluxation and then dislocation. Psoriatic arthropathy (see Fig. J.22) is often extremely destructive and, in addition, involves the distal interphalangeal joints. Small joint polyarthritis is also a complication of many medical diseases (see JOINTS, AFFECTIONS OF, p. 319). Gout (Fig. U.25) can be deceptive if the initial presentation is in the hand rather than the foot.
Osteoarthritis commonly affects the joints of the hand – in decreasing frequency, the distal interphalangeal joint, the proximal interphalangeal joint, the carpometacarpal joint of the thumb, the other carpometacarpal joints, and finally the intercarpal joints. The presenting symptoms are a gradual onset of stiffness, discomfort and weakness. Similarly, the physical signs slowly become more evident. Heberden's nodes are smooth, bony hard swellings situated dorsally on the radial and ulnar sides over the distal interphalangeal joint. They are due to osteophytes with overlying soft-tissue thickening. Occasionally, destruction of the joint may be more rapid. This is particularly seen in the proximal interphalangeal joint, and presents as a ‘spindle finger’. This must be differentiated from an acute ligamentous injury or early rheumatoid arthritis, which can produce similar physical signs. With very advanced finger osteoarthritis, there can be marked deformities with lateral or medial subluxation secondary to erosion of the articular surface. Bony crepitus can then be elicited on passive movement. All joints of the thumb can be involved. The classical deformity is fixed flexion of the trapezium-metacarpal joint, with hyperextension at the metacarpophalangeal joint and flexion at the interphalangeal joint (Fig. U.26). Isolated painful trapezium-metacarpal osteoarthritis is most commonly seen in middle-aged women and is accompanied by weakness; activities such as writing, knitting and unscrewing jars can be difficult.

**Tumours and tumour-like conditions**

*(See also LOWER LIMB, PAIN IN, 'Pain in and around the knee', p. 362; 'Tumours of bone', p. 371.)*

Almost all varieties of this complex pathological group can affect the hand. Fortunately, the most common bone lesions are benign. The enchondroma classically presents with a swelling of a phalanx, and on X-rays has a well-defined lytic lesion with a good margin and punctate areas of calcification. Unless complicated by a pathological fracture, they are usually asymptomatic. Lytic lesions in the terminal phalanx are most likely to be due to an epidermoid inclusion cyst, which is filled with keratin debris. The flexor tendon sheaths are a classical site for pigmented villonodular synovitis (giant-cell tumour of the tendon sheath). It presents either with a soft-tissue swelling on the palmar aspect of the finger along the tendon sheath or as a trigger finger. It can be locally aggressive. Beware of soft-tissue masses within the palm as these are most likely to be a malignant soft-tissue sarcoma. A biopsy is required to establish precise diagnosis.

**Miscellaneous**

Ganglia/mucous cysts are common in the hand, usually on the extensor surface overlying a joint (Fig. U.27). In the fingers, they are usually small and smooth, and have a hint of mobility and compressibility when examined with the finger extended. On flexion, they often feel bony hard. They may occur in association with an underlying osteophyte. On the palmar surface, they arise in conjunction with the fibrous flexor sheaths of tendons forming small, smooth, nodular swellings. Dislocations and sprains are common in the fingers. Dislocation of the terminal joint can easily be overlooked if the finger is examined a few hours after the injury due to the surrounding soft-tissue swelling.
Sprains (partial tears) of the collateral ligaments of the proximal inter-phalangeal joints are a cause of a ‘spindle finger’, with a tapered cylindrical swelling around the joint, which is often stiff and painful for several weeks. The X-ray is normal, in contradistinction to the spindle finger caused by osteoarthritis or rheumatoid. Gamekeeper’s thumb is due to a rupture of the ulnar collateral ligament of the metacarpophalangeal joint of the thumb. As with other complete ligamentous disruptions, this can be relatively painless and is easily missed unless the finger is correctly examined for instability; the thumb will collapse into abduction on making a pinch grip between the thumb and index finger. It is a common skiing injury.

URETHRA, FAECES PASSED THROUGH
Harold Ellis

Faeces or faecal fluid are passed per urethram only when the bladder is in fistulous communication with some part of the bowel, or with an abscess infected with *Escherichia coli* (see PNEUMATURIA, p. 519). The chief causes are as follows:

- Diverticular disease of the sigmoid colon with a fistula into the bladder (the commonest cause)
- Carcinoma of the bladder opening into the rectum, or into some loop of bowel that has become adherent to the bladder
- Carcinoma of the:
  - Rectum
  - Sigmoid colon
  - Caecum opening into the bladder either directly or through the medium of an intervening abscess
- Carcinoma of the uterus opening both into the bladder and into the rectum
- Crohn’s disease of the large or small bowel with vesical fistula
- Prostatitis or prostatic abscess opening into the rectum
- Rectovesical fistula from injury and sloughing, particularly after childbirth
- Appendiceal abscess opening into the bladder
- Pelvic actinomycosis

The passage of faeces into the urine may be simulated by some cases of very foetid cystitis due to infection by *E. coli*, especially in those who have diabetes.

If the symptom is due to carcinoma, it matters little which viscus is the primary site by the time the growth has involved both bladder and bowel. The differentiation resolves itself, therefore, between malignant and non-malignant conditions. If malignant disease is not obvious, it will nearly always be advisable to resort to surgical measures in the hope of discovering some curable primary condition – rectal, appendicular, prostatic or otherwise. This diagnosis will be suggested by the history and confirmed by local examination. Special investigations may include cystoscopy, barium enema and colonoscopy.

URETHRAL DISCHARGE
Ben Challacombe

Discharge of secretions or fluid from the urethra is usually only recognized in the male. The urethra is lined with accessory glands that normally produce small amounts of secretion. Urethral discharge is most commonly an accompaniment of sexually transmitted infections from *Neisseria gonorrhoeae, Chlamydia* or less often *Mycoplasma genitalium, Trichomonas vaginalis, Urea plasma urealyticum* or herpes simplex. In this setting, urethral discharge is commonly associated with DYSURIA (see p. 154), and it may also be complicated by epididymitis, orchitis or prostatitis. The diagnosis may be made by microbiological examination of appropriate samples of discharge fluid and urine.

Post-micturition dribble (PMD), which is the passage of a few drops of urine after micturition has been completed, is a normal finding in most males but becomes a problem for a few. It is due to pooling of urine in the bulbous urethra, and reassurance and education regarding manual perineal ‘milking’ out of urine is all that is usually necessary. A urethral diverticulum, which is rare in males, may be a cause of urethral discharge, if infected, or leakage of urine after micturition (PMD). After patch urethroplasty, the urethra may become baggy and dilated, and may be associated also with PMD. Infection within the Cowper’s glands in the proximal urethra may give rise to urethral discharge.

Females may develop urethral discharge, which usually comes from a urethral diverticulum. The discharge in females is usually purulent in nature as the discharge is infected.
URINE, ABNORMAL COLOUR OF
Mark Kinirons

The normal amber colour of urine is due mainly to urobilinogen; the depth of colour naturally varies with the concentration, and very dilute urine is nearly colourless. In very concentrated urine, the depth of colour may raise suspicions of biluria. Several substances can alter the colour of the urine. This is usually of no pathological significance, although a patient may seek an explanation; in a few conditions, the colour is characteristic and of diagnostic significance.

Bile pigment imparts a deep orange colour to the urine; in high concentration, the appearance resembles beer. This occurs mainly in bile outflow obstruction. Senna and rhubarb ingestion can produce a similar colour.

A red colour in the urine can be due to a number of substances. Haemoglobin is the most important, either in intact red cells, when the urine has a turbid or ‘smoky’ appearance, or as free pigment, when the urine is clear. This may be confirmed by a urine dipstick. If large amounts are present, and particularly if some of the haemoglobin has been oxidized to methaemoglobin, the colour may be brownish-black. In porphyric urine, the colour is typically that of ‘port wine’, but it may be pink or red. Myoglobinuria may give a red or brown colour in the urine. Other substances causing red urine include beetroot, blackberries, phenolphthalein in purgatives (if the urine is alkaline) and certain aniline dyes in sweets.

Apart from methaemoglobin, dark brown or black urine may be due to phenol (carboluria), melanin, homogentisic acid or p-hydroxyphenyl pyruvic acid. In carboluria, due to phenol poisoning, the urine may be greenish-brown. Melanin is found in the urine in some cases of disseminated malignant melanoma; the urine may be of normal colour when passed, but it turns black on standing, from above downwards. A similar colour change on standing occurs in the urine in alkaptonuria in which homogentisic acid is excreted. This substance also accumulates in the cartilage of the ear and in the sclera, which may become black, and in joint cartilage, causing severe arthritis; this syndrome is known as ochronosis. The urine may also darken in air in the very rare tyrosinosis in which p-hydroxyphenyl pyruvic acid is excreted. The antimicrobial drug metronidazole also causes the urine to become dark brown. Rifampicin also causes urine to go orange, and this may be used as a measure of compliance with therapy.

Due to the presence in normal urine of urobilinogen, any blue compound in low concentration may produce a green colour. Green and blue urines are most commonly due to biliverdin in long-standing obstructive jaundice, or to methylene blue in pills or sweets. Indigo-carmine and indigo-blue can also colour the urine. The former may rarely be present after exposure to industrial dyes, and the latter is the consequence of oxidation of indican. Indicanuria is due to intestinal malabsorption of tryptophan, which is metabolized to indole by intestinal bacteria in such conditions as coeliac disease and Hartnup’s disease.

URINE, INCONTINENCE OF
Ben Challacombe

Urinary incontinence is the complaint of any involuntary loss of urine. This is generally from the bladder and at times and in places that are inappropriate and inconvenient. Urine loss can be urethral or extrarethral secondary to anatomical abnormalities. Preservation of continence depends on the integrity of the lower urinary tract, both anatomically and physiologically. Incontinence secondary to an anatomical abnormality occurs congenitally, as in ectopic ureter, or is acquired, as in vesicovaginal fistula; physiological disturbance occurs because of an imbalance between the tone of the detrusor muscle and that of the external urethral sphincter.

Incontinence can be broadly divided into two main types – stress incontinence and urge incontinence – with both types far more common in women than men. Sphincter weakness results in genuine stress incontinence; in urge incontinence, detrusor activity is sufficiently enhanced to overcome the resistance offered by a normal sphincter mechanism. There are also several other rarer forms of incontinence, such as overflow incontinence, when the detrusor is flaccid so that urine trickles out when the fully distended bladder can hold no more.

STRESS INCONTINENCE

Mechanical damage to the sphincter is the most common cause of genuine stress incontinence. Broadly speaking, the sphincter apparatus in both sexes consists of three components: (i) the bladder neck, which is a muscle group derived from the detrusor muscle of the bladder wall (and the entrance to the prostate in men); (ii) the intrinsic urethral apparatus, which consists of muscle components from both the bladder neck and the external sphincter, together with fibrous and vascular components;
and (ii) the external sphincter, which consists of the striated muscle of the pelvic floor.

In women, all three components can be affected by stretch or direct damage during the passage of the fetal head during labour, giving rise to stress incontinence, particularly following multiple vaginal deliveries. This is due to urethral sphincter damage during childbirth, and to the weakening of the supporting pelvic floor muscles. The sphincter apparatus falls below the level at which it can be protected by transmitted pressure during coughing and other activity, so that the intra-abdominal pressure acts in an unopposed manner on the bladder dome. Increases in abdominal pressure produce a simultaneous leak of urine. The condition is commonly associated with anterior and posterior vaginal wall prolapses, manifested as cystocele and rectocele respectively, but this is by no means invariable. Stress incontinence can also develop due to a generally lax pelvic floor, obesity, chronic coughing, smoking and related medical conditions. Genuine stress incontinence also occurs in some congenital abnormalities, such as the short urethra, the wide urethra and epispadias.

Supporting the bladder neck by inserting the index and middle finger against the anterior vaginal wall and pushing upwards will control the leaking. This test mimics the effect of a successful surgical procedure, the options for which are currently transvaginal tape, transobturator tape or a colposuspension.

Significant stress incontinence is seen in up to 10–20 per cent of men following radical prostatectomy carried out for adenocarcinoma of the prostate. This is due to damage to the sphincter at the time of the operation during apical dissection and can be worsened by ajvant/salvage radiotherapy. It can be helped by pelvic floor (Kegel) exercises and training, but may require the insertion of a male sling or an artificial urinary sphincter. A period of some 6–12 months must elapse before the degree of remaining incontinence can be judged permanent.

External sphincter involvement by malignant extension from prostatic carcinoma is rarely enough in itself to give rise to stress incontinence, as the simultaneous obstruction produced by the enlarged malignant gland will compensate for loss of sphincter tone.

In men, a transurethral prostatectomy (TURP) or other bladder outflow procedures (HoLEP/PVP/open prostatectomy) for benign enlargement may rarely cause stress incontinence; the bladder neck is resected during the procedure, but there is occasionally additional damage to the external sphincter from poor technique, or in men with locally advanced prostate cancer where the usual landmarks are not easily visualized. More common following TURP is urge incontinence due to detrusor overactivity, which is often secondary to the initial bladder outflow obstruction.

The pelvic floor can be injured by trauma, such as gunshot wounds, and is particularly vulnerable to injury when the pelvis is fractured. A dual mechanism is often responsible for incontinence in the latter as direct damage to the pelvic floor is compounded by damage to its nerve supply, particularly the pudendal nerves.

**URGE INCONTINENCE**

Incontinence is called urge incontinence when the desire to micturate is so strong that it overrides all attempts of the sphincters to retain urine. Urge incontinence is more common in women than men but frequent in both sexes. The detrusor contracts in an abnormal manner during the filling phase and, as it does so, it opens the bladder neck and thus decreases the outflow resistance. This kind of incontinence is commonly due to idiopathic detrusor overactivity or neurogenic detrusor overactivity.

**Idiopathic detrusor overactivity**

This condition was formally known as bladder overactivity and the ‘irritable bladder’. It is an idiopathic condition and is often difficult to assess and frustrating to treat. The relationship between the symptom of urgency and urodynamic evidence of idiopathic detrusor overactivity is not always proven. Symptoms from involuntary detrusor contractions and detrusor weakness may be difficult to separate. Occasionally, involuntary detrusor contractions can be provoked by coughing, giving the impression of stress incontinence. Patients usually initially complain of frequency, urgency and nocturia, while urge incontinence occurs only in severe cases. Treatment involves behavioural modification with reductions in caffeinated drinks and bladder training, anticholinergic/antispasmodic drugs, botulinum toxin injection into the detrusor, neuromodulation and finally an augmentation ileocystoplasty in refractory cases.

**Neurogenic detrusor overactivity**

Incontinence related to neurological causes may be due to upper or lower motor neurone lesions. A variety of bladder and sphincter functioning states are possible. There may be abnormal bladder function, abnormal sphincter function or both. Either may be underactive or overactive, and all the various combinations of activity are possible.
In some upper motor neurone lesions, the central inhibitory impulses to the micturition centre in the sacral segments are lost. Sudden contractions of the detrusor muscle occur, resulting in unheralded micturition. The condition is associated with spinal cord injuries and multiple sclerosis, and follows cerebrovascular accidents. It is seen in some cases of Parkinson's disease, multisystem atrophy and syringomyelia, and it is a feature of normal-pressure hydrocephalus (Fig. U.28). The reflex centre in the cord is intact, so that the stretching of the bladder wall causes reflex detrusor spasm and micturition. A major difficulty is when an overactive sphincter generates high pressures during filling but fails to relax appropriately during voiding, and then the condition of detrusor sphincter dyssynergia arises. The result is bladder wall hypertrophy with trabeculation, formation of saccules and diverticula, and the presence of a urine residue. If the patient is paraplegic, management becomes extremely difficult, as sepsis, excoriation of the genitalia and perineum, and areas of pressure necrosis occur. In an upper motor neurone lesion, there is a significant risk of damage to the upper urinary tract due to high bladder pressures. Mechanisms to ensure detrusor pressure is minimized often need to be considered, for example regular intermittent self-catheterization, long-term catheterization and external sphincterotomy.

In many lower motor neurone lesions that affect the afferent and efferent portions of the sacral reflex arc as well as the reflex centre itself, the bladder is cut off from this spinal regulatory centre. As the pathognomonic feature of an upper motor neurone lesion is spasticity, so is flaccidity the feature of a lower motor neurone lesion. The detrusor muscle becomes flaccid, often insensitive to stretch, and the bladder distends enormously. The concomitant weakness of the sphincter mechanism eventually leads to overflow incontinence where urine trickles through the urethra. The bladder is readily palpable, asymmetrical, often enormous, not tender and relatively soft. Pressure on the bladder dome often results in the expression of urine, a diagnostic test which, in itself, can sometimes be adapted as a therapeutic technique to promote bladder emptying (Crede manoeuvre).

Lower motor neurone lesions are also associated with peripheral neuritis, as in diabetes. In diabetes, a selective peripheral neuropathy can affect bladder behaviour, as well as the mechanism of erection in the male, without any peripheral signs of such a neuropathy. Damage to the autonomic supply to the bladder may follow pelvic surgery, especially abdominoperineal excision of the rectum and radical hysterectomy.

Figure U.28 (a) Coronal MRI scan of normal-pressure hydrocephalus with massively dilated ventricles (arrowed). (b) Axial MRI of the same patient.
Fistulae from the urinary tract may communicate with the vagina, leakage may appear to be intermittent. If the abnormal opening is between the ureter and fistula, it is usually continuous but, incontinence from a hydronephrotic; it may drain only one calycine system. Be the ectopic ureter and the affected moiety is usually orthotopic ureter. The upper moiety ureter will always be heterotopic and opening inferior to the bladder, the ureter of the upper moiety is that the ureter of the lower moiety opens normally or into the urethra beyond the sphincter apparatus in small volumes, and normal controlled voiding may also occur with urine dripping from them. There is wide separation of the two pubic rami (pubic diastasis).

An ectopic ureter occasionally opens into the vagina or into the urethra beyond the sphincter apparatus in the female. Urine leaks continually, although usually in small volumes, and normal controlled voiding also occurs. The ectopic ureter usually drains the upper moiety of a duplex kidney but may be due to a complete ectopic kidney. The ectopic opening is often extremely difficult to find, but the intravenous injection of indigo-carmine or methylene blue will facilitate its localization. The Weigert–Meyer rule in duplex kidneys is that the ureter of the lower moiety opens normally into the bladder, the ureter of the upper moiety always being heterotopic and opening inferior to the orthotopic ureter. The upper moiety ureter will always be the ectopic ureter and the affected moiety is usually hydronephrotic; it may drain only one calycine system. Incontinence from a fistula is usually continuous but, if the abnormal opening is between the ureter and the vagina, leakage may appear to be intermittent. Fistulae from the urinary tract may communicate with the uterus (in which case urine can be seen escaping from the cervical os), or with the vagina (when the fistulae can usually be seen on speculum examination of the anterior vaginal wall). A ureterovaginal fistula may arise in the female from erosion of a calculus into one of the vaginal fornices, and also after gynaecological surgery. Vesicovaginal fistulae may follow hysterectomy, bowel resection, radiotherapy, cervical carcinoma, inflammatory bowel disease, pelvic fracture, childbirth and tuberculosis. In developing countries, the majority are associated with complicated childbirth due to pressure necrosis, and rarely with schistosomiasis. Investigation includes a cystogram, cystoscopy with bilateral retrograde studies, intravenous urogram or computed tomography urogram.

Rectovesical fistulae are due to diverticular disease, inflammatory bowel disease, complications of radiotherapy/HIFU/cryotherapy for prostate cancer, and malignancy of the bowel or bladder. They present with the classic symptom of pneumaturia, or with recurrent urinary infections or the passage of urine per rectum. They usually require surgical correction as diversion via catheterization is rarely successful.

The investigation and diagnosis of incontinence when there is no overt cause such as fistula or congenital abnormality depends on an accurate history and clinical examination. Specific investigations include a frequency–volume chart detailing voided volumes, incontinent episodes, fluid intake and degree of urgency, and a midstream urine specimen sent for microscopy and culture. Cystoscopy and urodynamic assessment of the patient may be required.

Urodynamics cystometry will measure the abdominal leak point pressure, which is the minimal pressure at which leakage occurs on straining. Video urodynamics indirectly look at the bladder neck and urethra and may give additional information on anatomical or neurological abnormalities in complex cases. Detrusor pressure is measured by subtracting the abdominal pressure from the total bladder pressure via pressure transducers in the rectum and bladder.

The bladder is usually a perfectly compliant organ in that intrinsic bladder pressure does not rise as the bladder fills. The normal curve is observed in genuine stress incontinence. In idiopathic/neurogenic detrusor overactivity, an uninhibited pattern of behaviour is seen. The ‘unstable detrusor’ is manifest as waves of contraction of a pressure that exceeds 10 cmH₂O. In lower motor neuron lesions, or in bladders affected by chronic retention, filling goes on and on with little alteration in intrinsic bladder pressure.
During the voiding phase, the bladder voiding pressure and the flow rate are measured. When there is sphincter incompetence, the voided pressure is low, while flow rate is often abnormally high; in obstruction, the voiding pressure will be high and the flow rate low.

**URINE, RETENTION OF**

Ben Challacombe

Retention of urine is the inability to empty the bladder completely; the end result is the acute, acute-on-chronic or chronic accumulation of urine within the bladder. In **acute retention**, there is a sudden complete inability to pass urine. The condition is painful and presents as a urological emergency. In chronic retention, there is a gradual increase in bladder size, often reaching enormous proportions of several litres (Fig. U.29). Pressure effects on the upper tract are not uncommon, and renal failure, in association with bilateral hydronephrosis and hydroureters, may be seen. Retention of the urine must be distinguished from anuria, when the kidneys fail to produce urine. In retention, whether acute or chronic, the kidneys still function, and urine continues to collect in the distended bladder. Urinary retention can be spontaneous or precipitated by another cause. Those with the latter may need no further intervention if the cause can be identified and treated, while those with spontaneous retention may require pharmacological or surgical treatment. Alpha-blockers and 5-alpha-reductase inhibitors are both successfully used in the treatment of lower urinary tract symptoms after a successful trial of voiding following acute urinary retention.

**ACUTE RETENTION**

Acute retention produces severe pain. The bladder is often palpable, central, tense, tender and dull to percussion. The most common cause in the male is outflow tract obstruction, secondary to **benign prostatic enlargement**. It should be noted that the severity of outflow tract obstruction bears no relation to the size of the prostate itself, as a tiny prostate can often be responsible for an acute retentive episode, and a huge prostate may remain asymptomatic. After urgent catheterization, it is vital to record the volume drained as this provides important prognostic information to guide management. In acute retention, catheterization produces a residual volume of between 500 ml and 1000 ml. Less than this and the diagnosis itself is in doubt; more, and chronic retention is more likely.

Common causes of acute urinary retention are:

- Benign prostatic hyperplasia
- Bladder neck stenosis/high bladder neck
- Prostatic adenocarcinoma
- Urethral stricture
- Tight phimosis or meatal stenosis
- Haematuria causing clot retention
- Urinary tract infection: cystitis or prostatitis
- Inguinal-scrotal and abdominal surgery
- Post-urological procedures: e.g. cystoscopy, prostatic biopsy, transurethral resection of bladder tumour
- Impacted urethral calculus
- Pelvic fracture
- General and spinal anaesthesia
- Immobility after other surgical procedures: hip/knee replacement
- Neurological: spinal cord injury, prolapsed intervertebral disc, multiple sclerosis, diabetic cystopathy and herpes virus

Acute retention may also be precipitated by overindulgence in **alcohol**, and the administration of **anticholinergic agents** for storage lower urinary tract symptoms, or **bronchodilating agents** (ephedrine) often used as cold remedies.

Acute retention of urine secondary to urethral intraluminal causes is most commonly the result of the **impaction of a calculus** in the urethra, which may be felt on examination of the penis or at the external urethral meatus. The history in such a case is dramatic, in that a normal flow of micturition is quite suddenly interrupted, causing a sudden pain along the urethra and the dribbling of a few drops of blood. **Blood clot** may impact in the urethra (‘clot retention’) as a complication of bleeding from any cause along...

**Figure U.29** Distension of the bladder to the umbilicus in chronic retention of urine due to benign prostatic hypertrophy.
URINE, RETENTION OF

the urinary tract, for example a renal carcinoma, or following any operation on the urinary system. Consequently, clot retention is not an uncommon complication following transurethral prostatectomy and a resected chip of prostate may also cause an obstruction.

Blockage of the bladder neck by the free-floating area of a pedunculated bladder tumour is rare; the growth is forced into the orifice during micturition, causing obstruction.

Complete traumatic rupture of the urethra leads to acute retention of urine. It almost always follows fracture of the pelvis but may follow a blow to the perineum, for example from a boot or from falling astride a bar. There will be history of injury, and blood will appear at the external urethral meatus; perineal haematoma may not be evident in pelvic fracture but will certainly indicate local urethral trauma.

If urethral rupture is incomplete, the patient should be encouraged not to pass urine as extravasation may occur and cause a haematourinoma in the perineum. An ascending urethrogram may delineate the injury, and suprapubic catheterization may be required.

Prolapse of an intervertebral disc or a spinal tumour or metastatic deposit will occasionally cause acute neurological urinary retention from direct pressure on the nerve roots. Retention may occasionally be the presenting feature of the condition, which, from the spinal point of view, remains asymptomatic. Acute, but painless, retention of urine will invariably occur after traumatic transection of the spinal cord.

Acute retention is a fairly common complication of operations on the rectum and neighbouring organs. It may also follow operations on the hip (in both sexes) and hernia repairs.

Acute retention of urine may be a manifestation of psychological illness; an organic cause should be sought and excluded before any psychogenic element is ascribed to the diagnosis. Retention due to psychiatric illness is more usually a complication of therapy as tricyclic antidepressants are powerful anticholinergic agents that suppress detrusor activity to the extent that retention, either acute or chronic, may occur.

Acute retention in female children is unusual, but it not infrequently occurs in males. While the male infant is still wearing nappies (diapers), ammoniacal ulceration of the foreskin, or of the meatus if circumcised, can give rise to acute retention because micturition is so painful that the child refuses to allow the act to continue. Retention in the presence of a tight phimosis is unusual, but the pathognomonic feature will be ballooning of the foreskin during attempted micturition. Meatal stenosis following ulceration secondary to circumcision can occasionally lead to acute retention in young males; when micturition is attempted, the urethra can be often felt as a distended and rigid band on the ventrum of the penis. Retention may also follow the inadvertent insertion of a foreign body into the anterior urethra.

Acute retention in women is relatively rare, and is commonly caused by urinary tract infection, prolapse and significant pelvic masses; it is sometimes associated with pregnancy, especially when the gravid uterus is retroverted, and large uterine fibroids may produce the same effect. Rarer causes include urethral diverticula, Fowler’s syndrome (impaired relaxation of the external urinary sphincter and polycystic ovary syndrome), post-surgery for stress incontinence, and urethral stricture. After urethral catheterization, a bimanual examination of the pelvis is mandatory, urine should be sent for culture, and a pelvic ultrasound examination should be ordered.

In young women, herpes genitalis, acquired as a sexually transmitted disease, may also precipitate acute retention, not only because of the urethral oedema associated with the herpetic lesions, but also by neurological involvement of the sacral reflex arc in such a way that detrusor activity is lost. Herpes zoster may give rise to acute retention in both sexes; in this case, the pathognomonic herpetic eruptions will be seen over the buttock area or the sacrum.

CHRONIC RETENTION

Chronic retention is insidious in its onset, and symptoms develop so slowly that the patient may deny significant lower urinary tract symptoms. It is only in retrospect and after surgical correction that the patient admits that there were significant lower urinary track symptoms preoperatively. Nocturnal enuresis (bed-wetting) is a classical presenting feature of overflow incontinence due to chronic retention. Many men present due to an acute precipitating episode on the background of chronic retention with a residual volume above 1–1.5 litres. An important subgroup is men with high-pressure chronic retention (HPCR). These men continue to void with a bladder volume above 800 ml and an intravesical detrusor pressure in excess of 30 cm H₂O. They have hydroureretonephrosis and, untreated, will develop renal failure. Abdominal examination reveals an enlarged tense bladder that is dull to percussion. After catheterization, intravenous fluid replacement with normal saline may be required due to a marked post-obstructive diuresis that
causes postural hypotension and dehydration. These men require long-term catheterization, intermittent self-catheterization or surgery for bladder outflow obstruction, as catheter removal always results in recurrent HPCR, with dangerous consequences.

*Benign prostatic enlargement/hyperplasia (BPE/BPH)* is unusual below the age of 50, but bladder neck stenosis or a high bladder neck can occur much earlier. On digital rectal examination, benign prostatic enlargement may be found. The gland may be smooth, uniform in consistency, elastic and movable within the pelvis when the enlargement is benign. A nodular, hard, irregular prostate may indicate *carcinoma*, while fixation to either side of the pelvic walls implies malignant extension well beyond the confines of the capsule. With bladder neck stenosis/hypertrophy, the prostate is of normal or small size.

When retention of urine follows *acute prostatitis* or a prostatic abscess, a history of recent urethral discharge will be obtained. The patient will be obviously ill, with a fever, rigors, frequency of micturition, pain on micturition, and perineal and pelvic discomfort.

Retention secondary to urethral *stricture* can be related to previous episodes of sexually transmitted diseases, commonly *Chlamydia trachomatis* or primary infection with *Neisseria gonorrhoeae*. Other common causes are urethral trauma, balanitis xerotica obliterans, and previous catheterization or instrumentation of the urinary tract. The history in stricture is of a gradual increasing difficulty with micturition, slowing of the stream, and dribbling after micturition; this last symptom occurs because of the column of urine that is trapped between the sphincter apparatus and the stricture. The investigation of choice is ascending urethrography, and subsequent urethroplasty when the length and position have been fully assessed. If instrumentation is undertaken as a primary investigation, the stricture may be carefully visually assessed before being divided via optical urethrotomy.
VAGINA AND UTERUS, PROLAPSE OF

Tony Hollingworth

A prolapse is the protrusion of an organ or structure beyond its normal anatomical position. Pelvic organ prolapse as defined by the International Continence Society is the descent of one or more of the vaginal segments: the anterior, the posterior, the apex of the vagina or, after hysterectomy, the vaginal vault.

The pelvic organs are supported by the pelvic floor, which in turn is made up of muscle, fascia and ligamentous support. The pelvic floor is made up from the levator ani, internal obturator and piriformis muscles, as well as the superficial and deep perineal muscles. The vagina is normally held in place by the transverse cervical ligaments (of Mackenrodt), the pubocervical ligament (pubocervical fascia) and the uterosacral ligaments. These vaginal supports have been divided into three levels by DeLancey. Level 1 is provided by the transverse cervical (Mackenrodt’s or cardinal) and uterosacral ligaments, level 2 by the anterior vaginal wall and rectovaginal fascia as the lateral attachment, and level 3 by the perineal body and perineal membrane as the distal attachment.

If the patient is examined in the left lateral or Sims’ position, with a Sims’ speculum holding back first the posterior and then the anterior vaginal wall, it is possible to determine which part of the vagina is prolapsing. This is classified into:

- **Cystocele:** prolapse of the deeper part of the anterior vaginal wall including the bladder
- **Urethrocele:** prolapse of the lower part into the anterior vaginal wall carrying the urethra in it
- **Cysto-urethrocele:** the combination of prolapse of both the upper and lower parts of the anterior vaginal wall, including the bladder and urethra
- **Rectocele:** prolapse of the lower part of the vagina including the rectum
- **Enterocele:** prolapse of the posterior fornix with the upper part of the vagina; this is related to the pouch of Douglas and may contain loops of small intestine
- **Uterine prolapse:** descent of the cervix with the uterus through the introitus, classified into four degrees:
  - First degree: the cervix descends to the vulva, but there is no protrusion through the introitus.
  - Second degree: the cervix protrudes through the vulva.
  - Third degree: the cervix is below the vulva.

- **Fourth degree (procidentia):** the whole of the uterus is outside the vulva (Fig. V.1).
- **Vault prolapse:** which can occur after hysterectomy, when the vaginal vault descends.

The POP-Q (Pelvic Organ Prolapse Quantification) classification of genital prolapse has been described to have a more objective, site-specific system for describing, quantifying and staging pelvic support. It measures nine sites on the vagina and perineal body in relation to the hymen and is useful in planning management (Bump et al. 1996The standardization of terminology of female pelvic organ prolapse and pelvic floor dysfunction, *AMJOG* 175:10–17).

Predisposing factors for the development of prolapse include weakening of the support mechanisms. The most common of these is injury as a consequence of childbirth and pregnancy; atrophy of the supporting tissues may also be implicated. Prolapse does occur in the nulliparous woman, but it is an unusual occurrence, and this suggests congenital or developmental weakness of the support structures.

There are a certain number of activating factors, which will increase the chances of prolapse development. These include obesity, an increase in intra-abdominal pressure in the form of chronic cough, constipation requiring continued straining or increased weight of the uterus, and if any undue traction has been put on the cervix at any time (Box V.1).

Symptomatology is related to the anatomical aspect of the prolapse (Box V.2), and many women complain of ‘something coming down’ or sitting on an egg. Although it is not painful as such, the patient may complain of backache. The woman may have urinary symptoms in the form of urgency. There may be difficulty in emptying the bowel, which requires

![Figure V.1 Fourth degree uterine prolapse procidentia.](image-url)
the woman to empty it digitally. There may be some discharge, and even bleeding and ulceration with a complete procidentia.

The differential diagnosis includes tumours of the vulva, vagina and cervix, which are not common tumours in the UK, hypertrophy of the cervix, a urethral diverticulum and true uterine inversion, which again is a very uncommon event. Fibroid polyp, chronic inversion of the uterus, vaginal cyst and endometrial polyp complete the list (Box V.3).

Management of these weaknesses can be either conservative or surgical, with the ring pessary being used in a number of patients while they await surgery, or if they are unfit or unsuitable for surgery. Surgery addresses the problem with either a suprapubic approach, particularly if there is any element of genuine stress incontinence, or a vaginal one. The primary surgery is vaginal hysterectomy and an appropriate repair. There has been an increasing use of meshes for these defects, but it is felt that these should be for a secondary repair rather than as a primary procedure. A complete procidentia would need to be reduced to prevent ulceration. It can be kept reduced by packing the vagina with a pack and local oestrogen cream prior to surgery, which would be the preferred option unless there is a contraindication to this.

VAGINA, DISCHARGE FROM

Tony Hollingworth

Discharge from the vagina can be classified in the following way:

- Physiological:
  - Varies with age and the time of the menstrual cycle
  - Common and may be the diagnosis when all else has been excluded

- Pathological:
  - Prepubertal
  - During reproductive life
  - Postmenopausal

PHYSIOLOGICAL DISCHARGE

The normal discharge from the vagina is a mixture of secretions from the uterine body, cervix and vaginal wall, the bulk of which originates from the cervix (Fig. V.2), as there are more glands there than in any other part of the lower genital tract. There are no glands in the vagina, since it is not a mucosa as such but a stratified squamous epithelium. The secretions vary during the menstrual cycle, being abundant, clear and almost free from leucocytes at the time of ovulation. At this time, the elasticity of the mucus is greatest (Spinnbarkeit), which allows easier penetration by the spermatozoa. At other times of the month, the cervical mucus is scanty, opaque and tenacious. The secretion from Bartholin’s glands, thin and mucoid, may be copious under sexual excitement, but under normal conditions it is scanty, so does not

Box V.1 Causes of prolapse

- Pregnancy and childbirth trauma
- Ageing and oestrogen withdrawal
- Smoking and chronic cough
- Constipation and straining
- Obesity
- Pelvic tumours
- Raised intra-abdominal pressure
- Previous surgery
  - Pelvic surgery
  - Bladder neck suspension
  - Burch's colposuspension

Box V.2 Symptoms of prolapse

- Feeling of discomfort
- Heaviness in the pelvis
- Lump coming through the introitus
- Worsening symptoms at the end of the day
- Dyspareunia
- Difficulty in inserting tampons
- Chronic lower back pain
- Urinary symptoms, urgency, frequency and incontinence
- Bowel symptoms difficulty or incomplete emptying of bowels

Box V.3 Differential diagnosis of a lump at the introitus

- Uterine prolapse
- Vaginal prolapse
- Cystocele
- Urethrocele
- Rectocele
- Enterocele
- Vault prolapse
- Vaginal cyst
- Imperforate hymen and haematocolpos (young women)
- Hypertrophy of the cervix
- Urethral diverticulum
- Cervical polyp or endometrial polyp (usually fibroid type polyp)
- Tumours of the vulva, vagina and cervix

Figure V.2 Normal cervix.
contribute to a vaginal discharge. The vaginal mixed secretion is acid in reaction, owing to the presence of lactic acid produced by Döderlein’s bacillus from the glycogen in the basal cells of the vaginal epithelium. This bacillus is normally found in the vagina from puberty to the menopause. The pH of the vagina is 4.5, the vaginal acidity being a bar to vaginal infection; unmixed uterine secretion is alkaline.

Normally, the amount of mixed vaginal discharge should do no more than just moisten the vaginal orifice; it may be increased with the presence of an ectropion (Fig. V.3), where there is eversion of the columnar epithelium towards the vagina. If excessive, the ectropion may require cautery or cryotherapy. Girls before puberty and women after the menopause do not have the protection of an acid secretion in the vagina.

**PATHOLOGICAL DISCHARGE**

**Prepubertal**

The main causes of discharge in a prepubertal girl include:

- **Non-infective:**
  - Poor hygiene
  - Foreign body
  - Sarcoma botryoides – a rare tumour

- **Infective:**
  - Threadworms
  - Sexual abuse

In young girls presenting with vaginal discharge, the most common diagnosis is due to a foreign body. This may necessitate ultrasound scanning or an examination under anaesthetic (EUA). At the time of EUA, a small hysteroscope can be inserted into the vagina, and the irrigating fluid used may flush the foreign body out and so treat the problem.

Poor hygiene again is not uncommon, and appropriate advice can be given to the mother. Threadworms may give intense itching, especially at night. One needs to be cautious if sexual abuse is considered, and the paediatric lead for child protection should be consulted. In the UK each hospital is now required to have a named doctor for child protection following the Climbie report. Sarcoma botryoides is a rare tumour that may present with discharge or bleeding in young girls and would need referral to a cancer centre for further management.

**Reproductive age**

Vaginal discharge in the reproductive age group, if not physiological, is most commonly due to infection. Cervical polyps and malignancy may present with excess discharge.

**Causes:**

- **Non-infective:**
  - Cervical ectropion, which is the eversion of the cervical endocervical columnar epithelium towards the vagina
  - Cervical polyps covered by columnar (glandular) cells
  - Malignancy
  - Foreign bodies, e.g. retained tampons or swabs – removal resolves the problem
  - General dermatitis
  - Allergies

- **Infective:**
  - Non-sexually transmitted:
    - Candidiasis
    - Bacterial vaginosis
    - Streptococcal infections
    - Anaerobes
  - Sexually transmitted:
    - Trichomoniasis
    - Gonorrhoea
    - Chlamydia
    - Pelvic inflammatory disease

In all cases, a relevant history should be obtained followed by a physical examination including a speculum examination to take the relevant swabs.

**Candida**

This is a common infection in women, which is often overdiagnosed. It gives rise to white patches of candida on the vaginal walls and cervix (Fig. V.4). It causes itching, discomfort and redness. It may complicate diabetes and pregnancy, as well as the use of antibiotics or the combined oral contraceptive pill (OCP). A swab may be taken for recognition of...
the mycelium and spores of *Candida albicans* in stained smears and for culture. Treatment can be topical or systemic azoles.

**Bacterial vaginosis**
This is characterized by a copious whitish discharge, which may be offensive or have a fishy smell. *Gardnerella, Mycoplasma* and anaerobes cause it. The vaginal pH is above 5, and the vagina is not inflamed. Diagnosis can be made by the amine test, in which some of the discharge is placed on a slide and potassium hydroxide is added to it; it then gives off a typical smell. Treatment involves either metronidazole or local clindamycin cream.

**Trichomonas vaginalis**
Caused by a flagellate parasite, this produces a frothy purulent discharge, causing local pain and soreness and being extremely irritating to the external genitalia. The discharge is green or greenish-yellow with small bubbles of gas in it, and it has a characteristic odour. The protozoon can be identified on microscopy. The trichomonads live in the vagina in symbiosis with the micrococcus *Aerogenes alicigenes*, which forms the froth or bubbles so characteristic of the discharge. It is a Gram-negative organism and causes the vaginal walls to have a typical red stippled (strawberry) appearance. Treatment is usually with metronidazole.

**Neisseria gonorrhoeae**
Gonorrhoea causes the cervix to be red, swollen and oedematous, being bathed in pus. There is nothing characteristic of gonorrhoeal discharge visible to the naked eye. The detection of the gonococcus can alone decide the question. This is often a matter of difficulty, because it is only in the few days immediately after infection that the organism can be found in the discharge. In chronic cases, the gonococcus must be looked for in one of three places: in the interior of the cervical canal, in the urethra, or in discharge squeezed from the orifices of the Bartholin’s glands. Gram’s method stains the discharge, the organisms being Gram-negative diplococci. Appropriate antibiotic treatment depends on the organism’s sensitivity, which may vary from locality to locality.

**Chlamydia trachomatis**
This is the commonest sexually transmitted disease in the UK and is caused by an obligate intracellular parasite, which lives in the columnar cells in the endocervical canal. It may cause discharge, or may not give any symptoms at all. The cervix may appear congested, friable and bleeds easily. An endocervical swab is needed for culture. Treatment involves doxycycline and there can be significant consequences to the woman’s fertility if it is not treated adequately.

**Pelvic inflammatory disease**
This condition presents with bilateral lower abdominal pain, vaginal discharge, low-grade pyrexia, tachycardia, bilateral adnexal tenderness and cervical excitation. The organisms involved include *Chlamydia trachomatis, Neisseria gonorrhoeae, Mycoplasma hominis* and anaerobes. This condition usually occurs in young women who are sexually active, so this should be covered in the history-taking. It is treated with antibiotics according to local protocols and local drug sensitivities, and it is important to treat properly on the first occasion as successive bouts of pelvic inflammatory disease may lead to infertility, tubal pregnancy, chronic lower abdominal pain and menstrual problems. Contact tracing of a partner should ideally be undertaken for *Chlamydia, gonorrhoea* and pelvic inflammatory disease.

**Postmenopausal women**
The differential diagnosis includes:
- Atrophic changes
- Complications of long-term use of a vaginal pessaries
- Pyometra
- Fistulae
- Malignancy (uterine, cervical, vaginal and vulval)

In postmenopausal women, the amount of vaginal discharge produced is reduced unless she is taking hormone replacement therapy. If she develops vaginal discharge, especially offensive in nature, a malignancy needs to be excluded. She will not have the infections that occur during the reproductive age except for *Candida*. One would need to exclude either an endometrial or a cervical lesion. In elderly women, a foul discharge may come from the interior of the uterus, a pyometra, in which case pus can...
be made to flow from the os uteri by squeezing the uterus or passing a sound; this may be due to senile endometritis, or it can be associated with carcinoma of the uterine body or cervix.

Fistulae may develop as a late manifestation of malignant disease, although this is not common, and it can also occur in bowel tumours and Crohn's disease. The rest of the women will have atrophic changes and may present with postmenopausal bleeding rather than discharge. If the woman needs a ring pessary inserted, this should be changed regularly; otherwise, discharge may develop.

**VAGINA, SWELLING IN**

Tony Hollingworth

Generalized swelling and oedema within the vagina may occur due to infection, which may be primary, for example *Candida*, or secondary, such as infected herpetic lesions. Condylomata (warts) may occur with a frond-like surface. Biopsy may be useful before instituting treatment. Otherwise, there are few structures that present as swellings within the vagina. If they do occur, they may be incidental findings during pregnancy or when taking a cervical smear. They may cause discomfort on intercourse or tampon insertion or may be found on self-examination. Patients may present with a lump in the vagina, the vast majority of which will be due to some form of prolapse (see VAGINA AND UTERUS, PROLAPSE OF, p. 730).

**BENIGN SWELLINGS**

- *Imperforate hymen*. The hymen is the membrane that partially covers the external vaginal opening. In some cases, this may be a complete block or occlusion. The result is not usually identified until puberty when the girl presents with cyclical pain but no menses. The menstrual flow is trapped behind the hymen and the girl develops a swelling arising from behind the symphysis, and actually presents with retention of urine due to the increasing pressure within the vagina. An examination of the vulva with parting of the labia will reveal a membrane across the vagina which is bluish in colour due to the blood collecting behind it (Fig. V.5). Ultrasound may be useful to confirm the diagnosis before performing a cruciate incision across the membrane to release the blood.

- *Simple mesonephric (Gartner's) or paramesonephric cysts* may be seen high up in the vagina in the fornices (Fig. V.6). They are embryological remnants, which have failed to be obliterated. They may be small, asymptomatic and found incidentally on vaginal examination. Occasionally, they can grow and give some degree of dyspareunia. The characteristic position and cystic feel serve to differentiate them from the various types of
vaginal prolapse. They can be treated by excision or marsupialization if necessary.

- **Small implantation cysts** may be seen at the vaginal orifice posteriorly; they are small, and may follow operations on the perineum or lacerations at childbirth. They may cause dyspareunia, and occasionally the scarring from removal means that there is no improvement of the symptoms.

- Occasionally, an endometrioma may burrow through into the posterior vaginal fornix from the floor of the pouch of Douglas into the rectovaginal septum, forming nodular growths, which tend to bleed at the time of menstruation. This condition may be confused with a primary carcinoma of the vagina. Biopsy and microscopic examination will confirm the diagnosis. It can also cause dyspareunia and may require surgery to resolve it, in some cases involving a general surgeon.

- **Benign tumours**: sessile and pedunculated swellings arise in the vaginal wall which on histology are found to be a papilloma, fibroma or lipoma. They are uncommon, and excision may be necessary if they interfere with intercourse.

**MALIGNANT SWELLINGS**

As in any type of malignancy, there is always the possibility of primary or secondary tumours. Primary tumours of the vagina are rare, and management needs to be undertaken at a gynaecological cancer centre. By and large, the prognosis from these tumours is poor despite radical surgery, radiotherapy and chemotherapy. The types of tumour are as follows:

- **Squamous lesions**: the vast majority, these usually occur in the upper vagina.
- **Clear cell carcinomas**: these were thought at one time to be related to exposure to diethylstilboestrol while in utero.
- **Malignant melanomas**: these may present with bleeding rather than swelling. The prognosis is very poor.
- **Endodermal sinus tumour**: this is a very rare type of adenocarcinoma.
- **Rhabdomyosarcoma (sarcoma botryoides)**: this is a rare tumour in girls less than the age of 5 years. It usually presents as vaginal bleeding. It has a characteristic appearance, like a bunch of grapes, and microscopic section proves its nature.
- **Secondary tumours**: these usually originate from the local organs, namely cervix and uterus, although there have been reports of secondaries from primary tumours in the ovary, colon and kidney.

Diagnosis is made by biopsy or excision biopsy depending on the size of the lesion, followed by magnetic resonance imaging of the pelvis and referral to a gynaecological cancer centre.

**VEINS, VARICOSE ABDOMINAL**

*Harold Ellis*

The point at which distension of veins becomes varicosity is arbitrary; most conditions that produce undoubted varicosity of the veins of the abdominal wall in some cases merely dilate them in others. When this dilatation is considerable, it nearly always has much diagnostic significance, particularly if the direction of blood flow is reversed.

Veins, when dilated, may be clearly seen as such, but they are unduly visible owing to wasting of the subcutaneous fat. Alternatively, in rare cases, they may be simply varicose, like veins in the leg, owing to idiosyncrasy or hereditary predisposition. In neither of these cases, however, is the blood flow in them reversed. To test the direction of blood flow, part of a vein should be chosen where there are no side branches, and the blood expressed from it by means of two fingers pressed gently down on the vein close together, and then drawn apart while pressure over the vein is maintained by each. When a length of the distended vein has been emptied in this way, one of the two fingers is lifted, and the time taken by the vein in refilling is noted. The procedure is repeated, the other finger being lifted off this time. It is then generally easy to decide whether the vein fills from below upwards, or from above downwards. Normally, blood flows from above downwards in the veins of the lower two-thirds of the abdominal wall. When the blood flow is from below upwards, there is almost certainly an obstruction to the inferior vena cava, the blood that is unable to return finding a collateral circulation via the abdominal wall to the superior vena cava.

Obstruction to the inferior vena cava is due to one or other of three main groups of conditions, namely:

- **Great general increase in the intra-abdominal tension**, owing to such conditions as ascites, ovarian cyst, or great splenic or hepatic enlargement
- **Thrombosis** without external obstruction
- **Obstruction by local compression**, especially by secondary deposits in the retroperitoneal lymph nodes

When the obstruction of the inferior vena cava is due not to the vein itself being thrombosed or invaded by new growth, but to the **general intra-abdominal pressure** becoming so great that the vein is, so to speak, flattened out, the varicosity of the veins upon
It is said that cirrhosis of the liver leads to varicosity of the veins around the umbilicus – the caput medusae. Most cases of cirrhosis of the liver cause no distension of the superficial abdominal veins until the general intra-abdominal tension has been greatly increased by the tenseness of the ascites, which occurs late. Not even the telangiectases that occur so commonly in men past middle-age around the lower part of the chest, in a line with the attachment of the diaphragm, indicate cirrhosis; these are quite as common in cases of emphysema without cirrhosis.

In summary, varicosity of the superficial abdominal veins generally indicates either thrombosis of the inferior vena cava, secondary to direct spread of thrombosis up to it from veins in the pelvis or in the leg, or else obstruction of the vena cava by secondary malignant disease.

**VERTIGO**

Michael Gleeson

Vertigo is a symptom for which there is a multitude of causes (Box V.4). Misdiagnosis must be avoided at all cost, and every effort should be made to establish the correct diagnosis by careful documentation of the patient’s history and a thorough clinical examination, followed by the selection and interpretation of specific investigations. While the patient may find accurate description of their symptoms difficult or beyond their capability to express, the physician should be absolutely clear that vertigo arising from the vestibular system is a subjective sensation of movement, which is usually rotatory. Patients may say that they feel either themselves or their environment moving. General sensations of light-headedness, dizziness or instability are rarely the result of labyrinthine or vestibular nerve disorders, and in these cases alternative causes should be sought.

Associated auditory symptoms, hearing loss and tinnitus are important features to establish and to define, as they are strongly indicative of an otological lesion. Assessment of this deficit is pivotal in subsequent management decisions. Almost all otological conditions that cause vertigo do so by involvement of the inner ear structures. A labyrinthine fistula should be seriously considered in patients with otorrhoea and those who have previously undergone surgery for chronic infection or otosclerosis.

Hennebert’s sign (the fistula sign) should be sought in these patients by applying pressure to the external meatus; if present, horizontal nystagmus will be observed. Careful otoscopy together with microsuction...
will most likely reveal a cholesteatoma or attic defect. Failing that, a defect in the otic capsule will be seen on a computed tomography (CT) scan (Fig. V.7).

Ménière’s disease is caused by fluctuation in endolymph production or resorption that results in distension of the membranous labyrinth, endolymphatic hydrops. The condition is characterized by episodes of vertigo that last for more than a few minutes and often several hours. They are often accompanied by pallor, prostration and vomiting, and preceded by a feeling of fullness in the affected ear, together with tinnitus that steadily increases in intensity. Marked nystagmus develops during the attack. As the vertigo subsides, the patient becomes aware of deafness in the affected ear, which over the course of several hours or days slowly resolves.

**Box V.4 Causes of vertigo and dizziness**

<table>
<thead>
<tr>
<th>General systemic</th>
<th>Haematological</th>
<th>Anaemia</th>
<th>Hyperviscosity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cardiovascular</td>
<td>Postural hypotension</td>
<td>Carotid sinus syndrome</td>
<td>Dyshytmias</td>
</tr>
<tr>
<td>Metabolic</td>
<td>Hypoglycaemia</td>
<td>Hyperventilation</td>
<td></td>
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<tr>
<td>Neurological</td>
<td>Supratentorial</td>
<td>Epilepsy</td>
<td>Syncope</td>
</tr>
<tr>
<td>Infratentorial</td>
<td>Multiple sclerosis</td>
<td>Vertebrobasilar ischaemia</td>
<td>Infections (e.g. syphilis, tuberculosis, herpes zoster)</td>
</tr>
<tr>
<td>Otological</td>
<td>Ménière’s disease</td>
<td>Post-traumatic syndrome</td>
<td>Positional vertigo</td>
</tr>
<tr>
<td>Miscellaneous</td>
<td>Ocular disorders</td>
<td>Odontogenic</td>
<td>Orthopaedic (e.g. cervical spine disease)</td>
</tr>
</tbody>
</table>

**Figure V.7** Computed tomogram of the labyrinth, showing a defect in the lateral semicircular canal (arrowed) caused by an osteoma that had developed in the mastoid antrum.

Positional vertigo is diagnosed when the history indicates that changes in head position precipitate vertigo, or that it can be demonstrated clinically. It can be caused by either a peripheral or central lesion. If peripheral, the condition is commonly attributed to disease of the otolith organ, cupulolithiasis, and severe vertigo is experienced, usually accompanied by rotatory nystagmus, when the head is lowered so that the affected otolith organ is undermost. The vertigo and nystagmus fades quickly while the head is maintained in this position. Usually, if the manoeuvre is repeated, the response is diminished or absent. This condition may develop after head injury, after a viral infection or for no apparent reason. In these circumstances, this condition is referred to as benign paroxysmal positional vertigo. Positional vertigo may also develop in association with central lesions such as multiple sclerosis or cerebellar tumours, often metastases. In such cases, there is no adaptation or fatigue of the response. Instead, the vertigo is present and persists for as long as the head is held in the critical position.

In cervical vertigo (which should not be confused with positional vertigo), the vertigo is induced by turning the head in a particular way. It can develop in patients suffering from atheromatous stenosis of the internal carotid artery or a similar narrowing of the vertebral arteries. It is thought that movement of the head causes obstruction to the cerebral blood flow, and there is some evidence that the vertebral artery may also be compressed within its bony canal by osteophytes in those with severe cervical spondylosis. Unlike positional vertigo, it is a movement of the neck, and not the position of the head, that determines the symptoms and, in some, it is not vertigo that they perceive but light-headedness; a few suffer syncope.
VESTIGO

Vestibular neuronitis is thought to be a viral infection of the vestibular nerve. It is associated with intense vertigo, with which a patient may wake and which may be accompanied by vomiting unless they lie absolutely still. There is no associated deafness or tinnitus. The intense vertigo usually passes off within a few days, but there remains a liability to brief vertigo on head movement that may persist for weeks or even months. Bithermal caloric tests of vestibular function show absent or severely reduced responses on one side.

Less common than any of the above are tumours of the VIIIth nerve or space-occupying lesions of the cerebellopontine angle or petrous apex, for example vestibular schwannoma, meningioma, petrous apex cysts and cholesteatoma. The most common of these is vestibular schwannoma (acoustic neuroma). Most patients with vestibular schwannoma present with a progressive unilateral hearing loss associated with tinnitus and episodic vertigo or instability of gait. As the tumour enlarges, other cranial nerve deficits develop that include facial paraesthesia and anaesthesia, trigeminal neuralgia caused by distortion of the Vth cranial nerve, dysarthria, dysphagia and voice disturbance secondary to IXth, Xth and XIIth cranial nerve palsies. Eventually, the brainstem becomes so severely moulded and compressed that hydrocephalus ensues with increasing headache and becomes so severely moulded and compressed that hydrocephalus ensues with increasing headache and visual disturbance (Fig. V.8).

More central lesions in the medulla and lower pons also cause vertigo. The vertigo in these conditions may be paroxysmal or continuous but, generally, the diagnosis depends less upon the quality of the giddiness than on other symptoms and signs arising from simultaneous involvement of structures adjacent to the vestibular nuclei. There is seldom tinnitus or deafness in any of these conditions, but long tracts may be involved, and nystagmus is present even when the patient is not actually feeling vertiginous. Moreover, the nystagmus is different in that, in labyrinthine and vestibular nerve lesions, the quick component is always towards the affected side, whereas in central lesions it is towards the left when the patient looks to the left, and to the right when he or she looks to the right. The diseases that produce vertigo in this situation include multiple sclerosis, thrombosis of the posterior inferior cerebellar artery, stenosis of the basilar artery, brainstem tumours and syringobulbia. Cerebellar disease can give rise to vertigo, more especially if the lesion is acute, as in penetrating injuries and infarction but, generally, chronic lesions cause a sense of disequilibrium rather than a sense of movement.

Systemic disorders also cause vertigo. Hypertension is common, dysrhythmias less so, while vertigo as the side effect of drug therapy is not infrequent. Demyelination and even syphilis should also be considered, if only to be excluded. Light-headedness often confused with vertigo accompanies anaemia and hypotensive states.

It is the author’s practice to undertake a neuro- otological examination for all patients with vertigo. This initial examination includes otoscopy, audiometry, evaluation of the central nervous system, and measurement of blood pressure and pulse rate, together with auscultation of the carotid arteries.

Audiometric and vestibular tests are tailored to each individual patient’s needs. A pure-tone audiogram alone may be adequate for some with absolutely normal, symmetrical hearing who give a classic history of a non-otological condition, for example vertebrobasilar ischaemia. The addition of a caloric test might be sufficient investigation for a patient with symptoms and signs of a benign otological condition, for example post-traumatic vertigo. More sophisticated tests are indicated and essential in other patients, for example those with asymmetrical or fluctuating sensorineural hearing loss, and absolutely mandatory for those in whom surgery is considered. They should also have a battery of basic blood tests to include a full blood picture, indices, biochemical screen and syphilis serology. A magnetic resonance scan is almost always arranged to exclude an intracanalicular tumour, and a CT scan is performed if surgical intervention is deemed necessary.

Containment of expense in the investigation of patients with vertigo has always been a problem for the otologist. Healthcare resources are becoming increasingly limited.
each year, and attempts to force economies on clinicians are more frequent. The surgeon should continually remember the potential consequences for the patient of missing the diagnosis of an acoustic neuroma or other cerebellopontine angle lesion at an early stage. For some, this could mean a permanently paralysed face and a lifetime’s misery. We should not be shy or slow to remind our institutions that the consequence for them could be a very substantial claim for negligence that would be difficult to defend.

In the UK, incapacitating vertigo precludes driving and also many other activities that might place either the patient or others at risk of bodily harm. While surgical treatment is often successful, the patient should not resume driving until they have been free from spontaneous attacks of vertigo for a period of at least 3 months. Standards of medical fitness to hold Heavy Goods Vehicles and Public Service Vehicles licences are much more stringent. In this respect, a history of recurrent vertigo or significant unilateral deafness is more than sufficient to disbar the patient from obtaining or renewing their licence.

VISION, DEFECTS OF

David Werring & Mark Kinirons

(See EYE, BLINDNESS OF, p. 172.)

In diagnosing the cause of visual loss, it is helpful to consider the speed of onset and permanence of loss, and whether one or both eyes are involved (Box V.5). In addition, the pattern of visual field loss is helpful in localizing the lesion to a particular part of the visual pathway, and it is sometimes characteristic of particular disorders (Table V.1). Lesions anterior to the optic chiasm (e.g. retina or optic nerve) cause monocular field loss. Chiasmal lesions (e.g. pituitary tumours) cause binocular, non-homonymous field defects (classically bitemporal hemianopia). Retrochiasmal lesions (optic tract, optic radiation or occipital cortex) cause homonymous field defects with varying congruity. A focal defect in the visual field is called a scotoma. A central scotoma is rapidly noticed and reported, and it is due to disease of the central retina or optic nerve. A centrocaecal scotoma (between the fixation point and the physiological blind spot) also localizes the lesion to the retina or optic nerve, and this may be seen with Leber’s hereditary optic neuropathy or toxic amblyopia. An arcuate scotoma is characteristic of glaucoma but may also be seen in optic disc drusen, optic neuritis and ischaemic optic neuropathy. An altitudinal scotoma (respecting the horizontal meridian) is typical of anterior ischaemic optic neuropathy. Peripheral constriction of the visual field occurs in open-angle glaucoma, retinitis pigmentosa and hysterical visual loss. In glaucoma, there is also cupping of the optic disc. Central vision may remain good, even though the field of vision is extremely limited. In retinitis pigmentosa, there may be associated night blindness. Retinitis pigmentosa may occur in isolation or in the context of a more widespread neurodegenerative condition, for example abetalipoproteinaemia, Refsum disease or adrenoleucodystrophy (typical bone spicule retinopathy), or mitochondrial cytopathy (‘salt and pepper’ retinopathy). The mode of transmission is variable, and there may be a range of associated neurological features depending on the underlying cause.

Constriction of the field of vision may also occur in hysterical blindness, but it is often variable or manipulable. A distinct psychological trigger can sometimes be identified, which is presumed to be somehow converted into a physical symptom (hence conversion disorder). Before making this diagnosis, all evidence of visual pathway disease must be carefully excluded.

<table>
<thead>
<tr>
<th>Box V.5 Differential diagnosis of visual loss</th>
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<tbody>
<tr>
<td><strong>Transient monocular loss</strong></td>
</tr>
<tr>
<td>• Thromboembolism (usually from the carotid artery)</td>
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<tr>
<td>• Retinal migraine</td>
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<tr>
<td>• Anterior visual pathway hypoperfusion (hypotension, hyperviscosity)</td>
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<tr>
<td>• Ophthalmological disease (e.g. intermittent angle-closure glaucoma)</td>
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<tr>
<td>• Vasculitis</td>
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<tr>
<td>• Demyelination (Uhthoff’s phenomenon)</td>
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<tr>
<td>• Visual obscuration</td>
</tr>
<tr>
<td><strong>Transit monocular visual loss</strong></td>
</tr>
<tr>
<td>• Migraine</td>
</tr>
<tr>
<td>• Cerebral hypoperfusion</td>
</tr>
<tr>
<td>• Malignant hypertension (thromboembolism, large-vessel stenosis, hypertension, hyperviscosity)</td>
</tr>
<tr>
<td>• Occipital seizures</td>
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<tr>
<td>• Visual obscurations (papilloedema)</td>
</tr>
<tr>
<td><strong>Sudden monocular non-progressive visual loss</strong></td>
</tr>
<tr>
<td>• Central retinal arterial or venous occlusion/branch retinal arterial or venous occlusion</td>
</tr>
<tr>
<td>• Traumatic optic neuropathy</td>
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<tr>
<td>• Retinal detachment</td>
</tr>
<tr>
<td><strong>Sudden binocular non-progressive visual loss</strong></td>
</tr>
<tr>
<td>• Vitreous haemorrhage</td>
</tr>
<tr>
<td>• Anterior ischaemic optic neuropathy</td>
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<tr>
<td>• Hysteria</td>
</tr>
<tr>
<td>• Sudden binocular non-progressive visual loss</td>
</tr>
<tr>
<td>• Bilateral occipital infarction</td>
</tr>
<tr>
<td>• Pituitary apoplexy</td>
</tr>
<tr>
<td>• Leber’s hereditary optic neuropathy</td>
</tr>
<tr>
<td><strong>Progressive visual loss</strong></td>
</tr>
<tr>
<td>(monocular or binocular)</td>
</tr>
<tr>
<td>• Optic neuritis (demyelinating, infectious, granulomatous)</td>
</tr>
<tr>
<td>• Toxic amblyopia, drug-induced optic neuropathy</td>
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<tr>
<td>• Anterior visual pathway tumour (e.g. parasympomatic meningioma)</td>
</tr>
<tr>
<td>• Anterior visual pathway aneurysm (e.g. giant aneurysm of the supraclinoid carotid artery)</td>
</tr>
<tr>
<td>• Radionecrosis</td>
</tr>
<tr>
<td>• Druen</td>
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<tr>
<td>• Chronic papilloedema</td>
</tr>
<tr>
<td>(e.g. idiopathic intracranial hypertension or tumour)</td>
</tr>
<tr>
<td>• Cataract</td>
</tr>
<tr>
<td>• Macular degeneration</td>
</tr>
<tr>
<td>• Cancer-associated retinopathy</td>
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</tbody>
</table>

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Other localizing symptoms include metamorphopsia (distortion of straight edges), which is characteristic of retinal disease, and acquired loss of colour vision, which is characteristic of an optic nerve lesion.

**TRANSIENT MONOCULAR VISUAL LOSS**

The possible causes are listed in Box V.5. Of these, the most common manifestation is with classical amaurosis fugax, in which the patient reports a black curtain coming from above or below to extinguish vision either completely or in one hemi-field. However, amaurosis fugax – literally translated as ‘fleeting blindness’ – does not always conform to this pattern and can be described in many different ways. The visual loss typically lasts a few minutes, with recovery over 15–20 minutes (in the reverse direction to the onset). This presentation suggests an embolus passing across the retinal circulation, resulting in temporary occlusion of the central retinal artery (or a proximal branch) before dissolving and moving to the periphery. This implies an embolic source, and is most likely to be platelet embolism from the ipsilateral carotid or aorta (due to atheromatous disease). Amaurosis fugax may be a premonitory symptom of central retinal artery occlusion, which causes permanent monocular blindness. Amaurosis fugax requires prompt investigation, particularly of the ipsilateral carotid artery: the finding of a significant symptomatic carotid stenosis is an indication for urgent surgical or endovascular intervention. Amaurosis fugax is not always due to carotid thrombo-embolism; alternative possibilities include retinal migraine (assumed due to retinal vasospasm) or haemodynamic reductions in retinal perfusion.

**TRANSIENT BINOCULAR VISUAL LOSS**

Visual obscurations are fleeting, usually fading visual losses on performing the Valsalva manoeuvre, coughing or changing posture. This is an important symptom of raised intracranial pressure with papilloedema. Migrainous aura can usually be recognized by the time course (classically up to 30 minutes, but may be longer) with an evolving pattern of visual disturbance, often with typical positive phenomena (teichopsia), as well as visual loss in the form of hemianopic or scotomatous loss. Cerebral hypoperfusion in presyncope is characteristically a fading of vision over seconds, always on standing, and usually associated with a dulling of hearing, feeling of faintness and sometimes a ‘coat-hanger’ ache of the neck and shoulders, before the patient collapses. Recurrent episodes in particular circumstances may suggest a vasovagal mechanism (e.g. precipitation by emotional or unpleasant stimuli). Particularly in older patients, extracranial large-vessel arterial stenosis can also cause posturally induced cerebral hypoperfusion, with similar visual fading and other focal neurological deficits, but this is probably rare. Transient bilateral visual loss in isolation is not typical of vertebrobasilar transient ischaemic attacks (TIAs); in such TIAs, other posterior circulation symptoms are usually to be expected in addition to visual loss.
SUDDEN MONOCULAR NON-PROGRESSIVE VISUAL LOSS

The important causes of sudden monocular blindness include acute central retinal artery or vein occlusion, anterior ischaemic optic neuropathy (which may be secondary to giant cell arteritis), vitreous haemorrhage and retinal detachment. In embolic central retinal arterial occlusion, there may be warning episodes of amaurosis fugax. Other causes of central retinal artery occlusion include in situ occlusion due to extensive atheroma (often in older patients in the context of diabetes and/or hypertension) and vasculitis (giant-cell arteritis, polyarteritis nodosa or Raynaud’s disease). A waxing and waning or stuttering onset is particularly characteristic of anterior ischaemic optic neuropathy, which may be either arteritic (due to giant-cell arteritis) or non-arteritic (due to in situ atheroma). Papilloedema is usually seen due to optic disc infarction. It is important to recognize anterior ischaemic optic neuropathy due to giant cell arteritis, since urgent treatment with high-dose corticosteroids may save vision. Associated features include systemic malaise, fatigue, headache (with scalp tenderness), myalgia and jaw claudication, and on examination the temporal arteries may be pulseless and thickened. The erythrocyte sedimentation rate may be markedly elevated, sometimes above 100 mm per hour. Giant-cell arteritis is nearly always limited to patients over 60 years old. Posterior ischaemic optic neuropathy is less common, and it may result from prolonged or severe perioperative hypotension.

The features of vitreous haemorrhage or retinal detachment can be detected on detailed ocular examination, including fundoscopy.

SUDDEN BINOCULAR NON-PROGRESSIVE VISUAL LOSS

This is usually caused by vascular pathology, either pituitary apoplexy or bilateral occipitoparietal infarction. Pituitary apoplexy is haemorrhagic infarction and necrosis of the pituitary gland, usually in the context of a pituitary adenoma. The associated features of this dramatic condition may include sudden collapse, headache, meningism and ophthalmoplegia. Pituitary apoplexy can occur postpartum with no adenoma present (Sheehan’s syndrome). Bilateral occipital infarction may be due to thrombotic or embolic occlusion of the posterior cerebral arteries (or the distal basilar artery, from which they arise). It can also occur with border zone (watershed) ischaemia and infarction due to prolonged reduced cerebral perfusion pressure, usually in the context of persistent systemic hypotension, for example after cardiac arrest. The pupillary light reflexes are preserved. Anton’s syndrome (the combination of cortical blindness, denial of the visual defect and confabulation) may result from bilateral posterior cerebral infarction. Leber’s hereditary optic neuropathy is a mitochondrial disorder (maternal inheritance) that usually affects males in the second or third decade, and causes sudden central visual loss. Typically, one eye is affected, followed by the second eye after an interval of weeks to months, but simultaneous loss can occur.

PROGRESSIVE VISUAL LOSS

Optic neuritis

The most important cause of monocular progressive visual loss in patients under 50 years old is optic neuritis, which may either be ‘idiopathic’ or occur in association with multiple sclerosis. Visual loss progresses over hours to a few days and reaches its nadir within a week. The severity of loss ranges from impaired colour saturation or contrast sensitivity with normal acuity, to no perception of light. Pain is usually present on eye movement at onset or shortly afterwards; the pain does not usually persist beyond a few days. Recovery begins within a month, stabilizing by about 6 weeks, and in the vast majority is good (better than 6/9 acuity). Partial recovery occurs, but lack of recovery is rare. Papilloedema may be present if the anterior portion of the nerve is affected. If the optic neuritis affects the more posterior retrobulbar portion of the optic nerve, initial fundoscopy is normal. Optic atrophy subsequently develops but does not correlate with the extent of visual recovery. Atypical features such as age greater than 50, lack of pain or prolonged pain, and an absence of the typical natural history of recovery should alert the clinician to other causes, for example ischaemic optic neuropathy, ‘atypical’ optic neuritis (e.g. post-infectious, infectious, post-viral, granulomatous or vasculitic) or compression by tumour. Bilateral simultaneous or sequential optic neuritis may occur, particularly in childhood. Treatment with corticosteroids may speed up visual recovery (although not improve final outcome), and may be offered to patients with bilateral involvement, pre-existing poor vision in the fellow eye, or severe visual loss and severe pain.

Optic neuritis may be associated with multiple sclerosis; about 50 per cent of patients with isolated optic neuritis will subsequently develop multiple sclerosis, although the interval is very variable. The presence of typical lesions on magnetic resonance imaging (MRI) increases the likelihood of developing
multiple sclerosis, and the extent of MRI disease is predictive of future disability. Follow-up studies indicate that multiple sclerosis presenting with optic neuritis has a relatively benign prognosis. The association of optic neuritis (usually bilateral) and subsequent or preceding spinal cord inflammation is termed Devic’s neuromyelitis optica; there are usually few cerebral lesions, and no oligoclonal bands are found in the cerebrospinal fluid (in contrast to multiple sclerosis, in which lesions and oligoclonal bands are almost invariably detected).

**Toxic amblyopia (nutritional optic neuropathy)**
The patient reports dimness or blurring of near or distant vision progressing over days to weeks. There is typically a central loss of vision for colours – green only in the earlier stages, but subsequently green and red. On examination, there may be a central scotoma, larger for coloured than for white objects. The fundus may be normal, but pallor of the temporal portion of the optic disc may be seen in some cases. Even though it is often termed tobacco–alcohol amblyopia, there is strong evidence that it is due to nutritional deficiency, rather than toxicity from alcohol and tobacco use, although the precise deficiency is not known. B vitamins have been suggested; B12 deficiency can cause an optic neuropathy, but this does not fully explain the majority of cases.

**Drug-induced optic neuropathy**
Treatment with drugs including ethambutol, isoniazid, streptomycin, chloramphenicol and quinine can cause an optic neuropathy. In the case of ethambutol and isoniazid, the optic nerve damage is dose-dependent. A careful history of the duration and timing of exposure to drugs in relation to visual symptoms is needed to make an accurate diagnosis.

**Compressive lesions of anterior visual pathways**
The hallmark of visual loss due to compressive lesion is a slowly progressive onset without recovery (in contrast to typical optic neuritis). Other features that raise the possibility of a mass lesion include complete absence of pain on eye movement, persistent pain over more than a week or two, proptosis, or spread to involve the other eye. The most common compressive lesions are pituitary region tumours (adenoma or craniopharyngioma), meningiomas, gliomas and aneurysms. In any case of visual loss without a history of typical optic neuritis, urgent imaging of the anterior visual pathways with computed tomography or magnetic resonance imaging (if available) is mandatory. The other main causes of progressive visual loss are listed in the Box V.5 above.

**Chronic papilloedema**
Progressive visual loss may result from very long-standing chronic papilloedema due to raised intracranial pressure. This may be due to a mass lesion and hydrocephalus, meningeal disease or, if no underlying cause is identified, idiopathic intracranial hypertension. Visual acuity is usually well preserved despite marked papilloedema, as the pathology does not directly damage the optic nerve fibres until late on in the illness.

**Cancer-associated retinopathy**
This rare retinal degeneration is a paraneoplastic manifestation of a remote systemic carcinoma. The typical presenting complaint is of shimmering visual disturbance. Specialist retinal electrodiagnostics are required for an accurate diagnosis.

**Developmental amblyopia**
Binocular fusion of images from both eyes is normally achieved by the age of 6 months. If one eye receives a defective image, for example from an uncorrected high refractive error, the less clear image is suppressed. If uncorrected, the eye becomes amblyopic and cannot be corrected by refraction. Alternatively, an imbalance of extraocular muscles that would cause diplopia in adults leads to a failure of development of visual pathways in the affected eye (strabismic amblyopia). Severe amblyopia may result from obstruction of the light pathway to the retina from birth, for example from congenital cataract or ptosis (stimulus-deprivation amblyopia).

**VISION, SUBJECTIVE DISTURBANCES OF VISION, SUBJECTIVE DISTURBANCES OF**
David Werring & Mark Kinirons

The main categories of visual loss have been described under VISION, DEFECTS OF (p. 739). There are other visual disturbances that do not fit easily into the above described scheme, and some of these will now be briefly considered. They can be divided into positive symptoms (an abnormal ‘excess’ of an aspect of visual perception) or negative symptoms (an abnormal loss of an aspect of visual perception), or distortions of normal visual perception. Some of the main causes are listed in Box V.6.

**POSITIVE DISTURBANCES**
Subjective visual disturbances, positive more than negative, are common in migraine. There are a wide variety of reported visual symptoms, including flashing lights (photopsia), shimmering, and transient spots before the eyes (scotomas). Fortification spectra (teichopsia), comprising zigzag or other
Charles Bonnet syndrome is seen usually in elderly patients, who report vivid scenes of detailed patterns, often in tessellating patterns; figures in ornate period costume are also typical. The hallucinations typically last minutes only, and are often seen at twilight. They are benign and non-threatening. The mental state is normal, and the rest of the visual world is normally perceived. This syndrome is probably due to cortical mechanisms resulting from a defective visual input of any ocular or neurological cause (e.g. cataract or macular degeneration).

NEGATIVE DISTURBANCES
Disturbances of colour vision are a hallmark of optic nerve disease, but they can also be caused by cortical lesions (e.g. stroke, demyelination, tumour or degenerative disorders) of the ventral occipitotemporal region (area V4), in which case visual acuity will be preserved. An inability to recognize familiar faces (or to learn new ones) despite normal objective tests of visual acuity is termed ‘prosopagnosia’, and this may be caused by lesions in the nearby fusiform gyrus. **Visual agnosia** refers to an inability to identify objects despite normal visual acuity. This may be due to difficulty in perception (**apperceiptive agnosia**) or in linking a perception to semantic knowledge of an object (**associative agnosia**). It is due to a lesion in the visual association pathways. **Balint’s syndrome** is classically due to a lesion in the occipitoparietal region and comprises an inability to reach for objects accurately (optic ataxia, or visual disorientation), an inability to fixate on a target (psychic paralysis of gaze), and an inability to perceive a scene as a whole despite perceiving its individual elements (simultanagnosia). **Alexia** is an inability to read despite normal visual acuity and, when not associated with a more general language (aphasic disorder), is termed ‘alexia without agraphia’. The syndrome is due to stroke in the posterior cerebral artery territory (occipitotemporal cortex). Disturbance of motion perception (akinetopsia) has also been rarely reported with lateral occipitotemporal cortical lesions.

DISTORTIONS
**Metamorphopsia** (distortion of straight lines) is usually due to retinal disease. Distortions of the sizes of objects (**micropsia** or **macropsia**) may be a manifestation of occipital lobe seizures or migraine aura. Micropsia and macropsia may accompany distortions of body image; this phenomenon has been termed **Alice in Wonderland syndrome**, in reference to the illusions described in the book by Lewis Carroll (Charles Dodgson), himself a migraine sufferer.
Voice is created by the modulation of sound produced by vocal fold vibration as air is forced or injected through the larynx from the lungs. Pitch is determined by the length and tension or stiffness of the vibrating vocal fold. Loudness of voice is a character largely determined by the pressure of air presented to, or allowed to develop in, the larynx. The subtlety and complexity of vocal fold vibration has only been appreciated in recent years with the development and widespread use of optical rods and stroboscopic light in clinical practice. With these tools, it is possible to see that the vocal folds vibrate in both planes of space, vertical and horizontal. A mucosal wave is generated during the production of voice as the epithelium covering the fold is able to slip on the underlying muscle and cord. Modulation of the sound produced in the larynx takes place in the upper airway – the vocal tract – and the quality or character of voice produced is largely determined by the shape and competence of these structures.

Even from this relatively simplistic outline of voice production, it can be appreciated that there are a multitude of causes of abnormal voice that range from structural lesions of the airway to disorders that affect laryngeal biomechanics. There is no entirely satisfactory classification of these disorders, but, from a clinical standpoint, the broad division into ‘structural’ and ‘functional’ conditions has much to commend it (Box V.7). The major weakness of this system lies in the fact that the larynx can compensate for disability, and that some voice abnormalities caused by covert structural lesions may present and be misdiagnosed at the outset as functional disorders.

While structural lesions cause hoarseness, functional disorders may produce a variety of more subtle voice changes, some of which are characteristic of their cause. The trained ear can detect these and phoniatrists refer to them in terms of ‘jitter’, ‘shimmer’, breathiness’, etc., descriptors that indicate the stability of a particular sound or smoothness of transition from one frequency to another. These abnormalities can be documented by waveform and spectral analysis of vocal fold vibration and voice frequency.

**STRUCTURAL**

**Acute laryngitis**

The most common cause in adults is a viral infection that accompanies a common cold or upper respiratory tract infection. Rhino-, parainfluenza, respiratory syncytial and adenoviruses are the most frequently implicated. Secondary bacterial infection often takes place. Patients present with generalized malaise and complete or intermittent loss of voice and cough. Spontaneous recovery takes place in a few days, aided by steam inhalations, cough suppressants and voice rest. Some require antibiotics, and it goes without saying that those who smoke should not.

Laryngitis in children is known as *croup*, and it is a more serious condition than its adult counterpart. The oedema associated with the infection in childhood may cause significant narrowing of the airway and critical obstruction. In addition to loss or alteration of voice, the child may develop stridor with intercostal and supraclavicular recession as he or she struggles to breathe. In severe cases, infection can spread down the airway: *laryngotracheitis* and *laryngotracheobronchitis*. These children can be extremely unwell, and both conditions can reach life-threatening proportions.

Bacterial infection of the supraglottitis, known as *supraglottitis* in adults and *epiglottitis* in children, is fortunately becoming less common as a result of immunization programmes against *Haemophilus influenzae* infection. Although *H. influenzae* is the most common organism involved, *Streptococcus pneumoniae*, *Staphylococcus aureus* and haemolytic streptococci may also cause the condition. Supraglottitis, and particularly epiglottitis, presents as a rapidly progressive illness with fever, sore throat,
hoarseness or muffled voice and dyspnoea. The diagnosis should be made on the basis of the clinical history and the speed of progression of the illness, which can be frighteningly fast. Inspection of the larynx in the awake child in order to see the swollen epiglottis is dangerous as it may incite acute laryngeal spasm and respiratory arrest. A delay in management to obtain a lateral neck X-ray that will show the epiglottic swelling can also be critical: better to examine the child under anaesthesia with the ability to secure the airway by tube or tracheostomy. In adults, the infection is rarely as severe, but suppuration is more common, with the development of an epiglottic abscess. Nevertheless, the condition demands close observation, humidification of inspired air, antibiotics and occasionally steroids.

**Angioedema or allergic laryngitis** of the larynx can be life-threatening and can be precipitated by a number of substances that include food colourants and common drugs such as angiotensin-converting enzyme inhibitors and non-steroidal anti-inflammatory drugs. In some, the condition is inherited through an autosomal dominant gene that causes C1 esterase inhibitor deficiency. In an attack, patients lose their voice, experience difficulty swallowing their saliva and develop swelling of the lips, tongue and periorbital tissues.

**Diphtheria** of the larynx is very uncommon nowadays, but it is still seen in developing countries and individuals travelling from them. Caused by *Corynebacterium diphtheriae*, it usually affects young persons. It presents with a fever, progressively severe sore throat, loss of voice and increasing airway obstruction. An exudative inflammatory response can be seen in the throat as a thick, grey-green, plaque-like membrane over the fauces, pharynx and larynx that, if removed, causes bleeding of the underlying tissues. The diagnosis is made by the bacteriological examination of swabs, smears and cultures. Administration of diphtheria antitoxin and the establishment of a secure airway by tracheostomy are life-saving measures, as mortality is largely due to cardiac dysrhythmias, neuropathic effects of toxins and airway obstruction. The organism is still sensitive to penicillin.

**Chronic laryngitis**

Regardless of the cause, chronic inflammation of the larynx can result in the development of erythema and/or oedema over the vocal folds (*Reinke's oedema*), discrete polyps or more widespread **polypoid degeneration** of the laryngeal mucosa and granuloma formation (Fig. V.9). Smoking, vocal abuse such as excessive shouting, intubation injury, radiation, chronic bronchitis, emphysema and laryngopharyngeal reflux are most often implicated, and these are sometimes loosely grouped together as non-specific causes of chronic laryngitis. Chronic candidal infection may be associated with inhaled steroid preparations. All these patients have a persistent and long-standing voice disturbance with no associated swallowing problem or pain. Dyspnoea is extremely unusual. The mainstay of treatment in this condition is correction or avoidance of the precipitating cause, vocal hygiene (voice rest, steam inhalations and avoidance of dehydration), antireflux measures and speech therapy. In some, the inflammatory reaction will subside, the larynx and voice returning to normal. Unfortunately, in others it does not, and microsurgical procedures become necessary, for example stripping excessive redundant mucosa from the cords or removing polyps.

![Figure V.9](a) The normal appearance of the vocal cords. Both cords are pearly white and well defined. (b) Long-standing chronic laryngitis has resulted in the vocal cords becoming hyperplastic and granular in appearance.
A number of specific causes of chronic laryngitis are still seen, especially in developing countries and in immunocompromised patients. These include tuberculosis, syphilis, scleroma, leprosy, blastomycosis, histoplasmosis, sarcoidosis and Wegener’s granulomatosis. It is important to exclude these conditions in patients who fail to respond to conventional therapy, or who have an unusual history with progressive dyspnoea and malaise being prominent. Diagnosis is made by biopsy, smears and cultures from swabs in infective disorders. In Wegener’s granulomatosis, the laryngeal condition may be part of a multisystem disorder affecting all parts of the respiratory system and the kidneys. Raised serum anti-neutrophil cytoplasmic antibody (ANCA) levels and biopsies confirm the diagnosis.

Neoplasia

Papillomata are the most common benign laryngeal neoplasms, which tend to affect children rather than adults. Multiple lesions develop within the larynx and cause hoarseness and progressive dyspnoea. In some, complete remission takes place at puberty. In a few patients, the papillomata spread into the trachea and bronchi. Diagnosis is made by biopsy. Malignant transformation is most unusual but is occasionally seen in adults who smoke.

Dysplasia of the laryngeal epithelium may also cause a change in voice quality. Degrees of dysplasia from mild to severe and in situ carcinoma are recognized on the basis of histological criteria. On laryngoscopy, they appear as white plaques, sometimes with adjacent erythematous areas. Some, although not all, progress to frank malignancy.

The majority of laryngeal cancers are squamous cell carcinomas. Three distinct sites within the larynx are recognized, and tumours developing in them tend to present and behave in slightly different ways. At the outset, supraglottic tumours cause a sensation of a persistent lump in the throat that may be painful when swallowing, the pain often radiating to the ear. As the tumour grows and spreads onto the vocal cords, hoarseness develops, followed by progressive dyspnoea. Some patients experience minor haemoptyses. In contrast, tumours that develop on the vocal cord present with huskiness and progressive dyspnoea, only developing problems with swallowing at a late stage (Fig. V.11). Subglottic tumours are most uncommon and present with dyspnoea (Fig. V.12). Nodal metastases in the neck are found at presentation in up to 25 per cent of patients with supraglottic tumours and 5 per cent with glottic carcinomas. Laryngeal cancer develops in heavy smokers. Any middle-aged smoker who develops hoarseness must have the diagnosis of laryngeal cancer excluded without delay by laryngoscopy and biopsy of any suspicious lesion. Squamous cancers respond well to radiotherapy when small, but more advanced tumours require surgical treatment. As this is a smoking-related disease, 15–20 per cent will develop second primary neoplasms in the upper aerodigestive tract, either synchronously or metachronously.

In contrast, blunt injury to the neck may result in a fracture of the larynx. The voice abnormality in this situation is associated with subcutaneous emphysema that is palpable and visible on X-rays, painful dysphagia and, in severe cases, dyspnoea.

Trauma

In its simplest form, constant vocal abuse through shouting inflicts various degrees of hoarseness that can be reversible as nodules (Fig. V.10), Reinke's oedema and submucosal haemorrhages subside.

Figure V.10 Vocal nodules at the junction of the anterior and middle third of the vocal cords.
Early myxoedema can present as a subtle voice change, huskiness or restriction of vocal range. With increasing oedema deposited in the vocal folds, dyspnoea and critical airway obstruction can develop alongside more classical signs of hypothyroidism. Amyloid infiltration of the larynx may be seen in both the primary and secondary forms of the disease. Abnormalities of voice develop when the infiltrate involves the vocal cord and, in severe cases, dyspnoea from airway obstruction may develop. Diagnosis is made on the basis of biopsies and plasma immunophoresis.

FUNCTIONAL

Neurological

Vocal cord palsy may be unilateral or bilateral and is caused by lesions of the vagus or recurrent laryngeal nerves. Paralysis of the left vocal cord is more common than that of the right cord; the longer course of the left recurrent nerve accounts for this difference. Palsies are not uncommon in clinical practice, and they present with sudden voice change that may be relatively subtle or very obvious. The degree of abnormality depends on the size of the glottic chink that develops as a result of loss of vocal cord adduction. High lesions that affect both superior and recurrent laryngeal nerve function tend to be worse, and the voice change is associated with aspiration of fluids and a bovine cough. Over time, the larynx often compensates for unilateral paresis, and voice quality can return to normal without recovery of vocal cord movement.

Iatrogenic trauma acquired during thyroid, oesophageal, carotid, cardiac and skull base surgery accounts for about 40 per cent of palsies. Malignant infiltration of the nerves at the skull base by glomus tumours, in the neck from thyroid or oesophageal carcinomas, and in the mediastinum from bronchial carcinoma, metastases or lymphomas is responsible for about 25 per cent. No cause is apparent in 25 per cent, knife and bullet wounds produce 25 per cent, and various neuropathies account for the rest.

Muscle tension dysphonia

This is caused by disproportionate, ineffective or uncoordinated contraction of the intrinsic and extrinsic laryngeal musculature. In some, endoscopy will show incomplete closure of the vocal folds on phonation, with excessive contraction of the vocalis muscle causing adduction of the false cords – dysphonia plicae ventricularis. In contrast, others show ineffective glottic closure with bowing of the vocal folds, as if the folds were intrinsically weak or subject to fatigue – laryngeal myasthenia or presbylarynges. This is particularly common in the elderly. Various surgical procedures have been suggested for both these conditions, but speech therapy and vocal rehabilitation are helpful if not curative in most.

Psychological

Hysterical dysphonia or aphony is not common and can be extremely difficult to treat. It may result from emotional trauma or reflect emotional instability. The important issue in this condition is to recognize it and the fact that the patient may have a serious underlying psychiatric illness that requires treatment in its own right.

Miscellaneous

Early myxoedema can present as a subtle voice change, huskiness or restriction of vocal range. With increasing oedema deposited in the vocal folds, dyspnoea and critical airway obstruction can develop alongside more classical signs of hypothyroidism. Amyloid infiltration of the larynx may be seen in both the primary and secondary forms of the disease. Abnormalities of voice develop when the infiltrate involves the vocal cord and, in severe cases, dyspnoea from airway obstruction may develop. Diagnosis is made on the basis of biopsies and plasma immunophoresis.
VOMITING

Simon Anderson

Vomiting is the forceful ejection of material from the upper gastrointestinal tract. It may or may not be associated with nausea (see p. 447) and should be distinguished from regurgitation (p. 447) whereby food is expelled from the oesophagus before reaching the stomach. Obstruction or partial obstruction of the gut lumen immediately comes to mind on considering possible causes. Intra-abdominal pathologies, however, as well as metabolic, systemic, psychogenic and physiological causes, such as pregnancy, must be considered.

Vomiting is a significant feature of mechanical obstruction secondary to lumen narrowing caused by inflammation (peptic ulcer disease, Crohn’s disease or ischaemia), fibrosis (healing of inflammation and surgery), neoplasia (adenocarcinoma, lymphoma or polyps) or foreign bodies (bezoar, worm infestation or gallstone).

It is important to bear in mind that obstruction of the gastrointestinal tract at any level may result in vomiting. It may, consequently, be a feature of obstruction of either the small or large bowel. Certain conditions are common to both areas (Crohn’s disease, anastomotic stricture, adenocarcinoma, and ischaemic strictures). Some conditions, however, are limited or more likely to occur in the small bowel, or are more problematic should they occur in this area (adhesions, herniae, lymphoma, hamartomatous polyps, Henoch–Schönlein purpura, mesenteric adenitis, serosal secondary deposits, ulcerative jejunitis, neuroendocrine tumours, bezoar, foreign body or gallstone impaction).

Mechanical causes for vomiting include volvulus. Volvulus of the stomach is an uncommon surgical emergency, which presents with upper abdominal pain and ‘fruitless’ vomiting. Volvulus of the rectosigmoid colon is not uncommon in the elderly, and it tends to recur if not addressed by surgical or endoscopic fixation. Muscle hypertrophy of the pylorus (pyloric stenosis) is well described in infants, and it presents with projectile vomiting.

Vomiting is a feature of acute appendicitis, where it is preceded by pain localizing to the right lower quadrant. It is, additionally, a common feature of both biliary and renal colic.

Insufficient emptying of the stomach (gastroparesis) is either idiopathic or due to poorly controlled diabetes.

It is diagnosed with a nuclear medicine ⁹⁹m-Tc gastric emptying study.

Infective causes of vomiting include those directly involving the gastrointestinal tract such as gastroenteritis (e.g. *Staphylococcus aureus* and *Bacillus cereus*), or epidemic viruses (e.g. Norovirus, Norwalk virus), and those outside the gastrointestinal tract e.g. viral and bacterial meningitis and syphilis (tabes dorsalis).

The ‘morning sickness’ associated with the first trimester of pregnancy is normally self-limiting and mild. Persistent vomiting associated with weight loss, dehydration and acidosis is termed ‘hyperemesis gravidarum’ and is part of a wider spectrum of conditions, including fatty liver or pancreas. The presence of haemorrhagic retinitis is a poor sign.

Iatrogenic causes of vomiting, in addition to adhesions and anastomotic strictures, include post-vagotomy/oesophagectomy pyloric obstruction, medication (opiates, non-steroidal anti-inflammatory drugs, steroids, azathioprine, cisplatin/chemotherapy, and allergic reactions) and ileus, which may occur following any surgical operation and is not limited to intra-abdominal procedures alone.

Vomiting is a well-recognized feature of intracranial pathology (space-occupying lesions, meningitis, cerebrovascular accident, hydrocephalus and middle-ear disease). The presence of the vomiting centre and chemoreceptor trigger zone in the medulla oblongata and fourth ventricle, respectively, gives rise to vomiting being a feature of systemic disease such as sepsis, malaria, pyrexia and diabetic ketoacidosis. Psychiatric and behavioural disorders are commonly associated with recurrent vomiting.

The complications of vomiting include Mallory–Weiss tear of the oesophagus, suggested by vomiting followed by haematemesis, and rupture of the oesophagus (Boerhaave’s syndrome), suggested by vomiting with severe, acute retrosternal chest pain. Vomiting during childbirth leading to the aspiration of gastric acid is termed Mendelson’s syndrome. Persistent vomiting leads to the development of a hypokalaemic alkalosis due to loss of gastric acid. In cases of bulimia, repeated vomiting causes erosion of dental enamel.

The cause of a patient’s vomiting is often indicated by the clinical setting and other organ-specific symptoms or signs. In cases of mechanical obstruction, the interval between eating and vomiting gives some indication of the level of a mechanical obstruction.
The contents of the vomitus may give an indication of the cause. In particular, the presence of a small volume of fresh or altered ('coffee-grounds') blood can indicate a primary upper gastrointestinal cause (see Haematemesis, p. 238) or occur as a consequence of vomiting (Mallory-Weiss tear). Similarly, the presence of bile implies little. Projectile vomiting is classically associated with pyloric channel obstruction or raised intracranial pressure; it occurs also in those with large antral gastric ulcers with no physical obstruction. The demonstration of a succussion splash is as likely to be due to some degree of gastroparesis as outlet obstruction, to which it is traditionally ascribed.

INVESTIGATIONS
The investigation of vomiting aims to determine both the underlying cause and its consequences (metabolic disturbance or aspiration pneumonia). General blood tests (full blood count, urea/electrolytes, serum calcium/phosphate, amylase, liver blood tests and serum glucose) can be very useful in both of these respects. Likely upper gastrointestinal pathology is most easily accessed by endoscopy, although care must be taken to prevent aspiration. Colonoscopy can be attempted and may prove useful, but an oral bowel preparation will not be possible. X-ray studies (contrast meal and enema) can be used to identify obstruction and paresis, but these are best performed using water-soluble contrast rather than barium on account of the risks of aspiration or worsening lower-bowel obstruction. They are, however, less sensitive than their barium counterparts. In the absence of any other positive findings, nuclear gastric emptying studies can quantify gastroparesis. The main role of oesophageal motility studies is to identify conditions such as achalasia or oesophageal scleroderma. If no gastrointestinal or systemic cause can be elicited, it is important to exclude intracranial pathology through a computed tomography, or preferably magnetic resonance imaging, scan of the head. Lumbar puncture will be necessary in cases of syphilis.

VULVA, SWELLING OF

Tony Hollingworth

The differential diagnosis of vulval swellings includes not only tumours of the vulval structures and skin appendages, but also swellings that appear at the vulva as a result of the displacement of other structures, as in cases of uterine prolapse and cystocele (see VAGINA AND UTERUS, PROLAPSE OF, p. 730). Hernias into this region can occur, and no further discussion is included in this section. Inflammatory lesions and ulceration of the vulva may be accompanied by swelling of the vulva due to associated oedema. These conditions are considered under VULVA, ULCERATION OF (p. 752). Conditions presenting with itching of the vulva as the main complaint are described under PRURITUS VULVAE (p. 529).

INFECTIVE SWELLINGS
- Warts (condyloma acuminata)
- Bartholin's abscess

Warts on the vulva are usually multiple (Fig. V.13). They are caused by the human papilloma virus (DNA virus) types 6 and 11 and are almost invariably transmitted sexually. They may spread throughout the lower genital tract and anal region. They have been associated with premalignant disease of the cervix. Vulval warts may proliferate and coalesce, in which case they are referred to as condylomata acuminata. This situation can be problematic in pregnancy and in patients who are immunocompromised, for example those with HIV or systemic lupus erythematosus, and those on long-term steroids.

Bartholin's abscess presents as an extremely painful swelling in the region of the Bartholin's glands and occurs at the entrance to the vagina. Pressure in the gland causes much pain, and the area appears reddened. The duct of the gland has become blocked and the secretions within the gland infected. It may

Figure V.13 Vulval warts.
discharge by itself, but treatment is usually surgical in the form of marsupialization to create a new duct or introduction of a balloon to open it. Recurrence may occur.

**CYSTIC SWELLINGS**

- Bartholin's cyst
- Sebaceous cyst
- Sweat glands
- Mucous cyst
- Implantation cyst
- Dermoid cyst
- Gartner’s duct (mesonephric) cyst

The common cystic swelling is a *Bartholin’s cyst* (Fig. V.14). This is usually occurs in the duct of the Bartholin's gland, producing a swelling in the posterior third of the labium majus that projects medially so as to encroach on the vaginal entrance, causing dyspareunia. It is not particularly tender unless it becomes infected, forming an abscess. The cyst tends gradually to increase in size, causing local discomfort, and treatment is as above.

Sebaceous cysts are fairly common, as a rule affecting the labia majora. They may occur in groups. Mucous, inclusion (Fig. V.15) and implantation cysts also occur, as do vestigial cysts of mesonephric origin (Gartner’s duct cysts). The true nature of these cysts is not usually known without histological examination.

The sweat glands in the vulva can become inflamed and produce a purulent discharge. These glands can coalesce and lead to hidradenitis suppuritiva which can lead to pain, swelling and discharge in the vulva.

**BLOOD CYSTS**

- Varicocele
- Traumatic haematoma
- Endometrioma

*Varicocele* of the vulva occurs mainly in pregnancy and can become worse with successive pregnancies. It gives a typical varicose appearance in the labia majora, and the woman can become conscious of an uncomfortable swelling on standing. The veins seldom rupture during delivery. Varicocele must be differentiated from an inguinal hernia extending into the labium majus and from a cyst of the canal of Nuck (a processus vaginalis that has failed to become completely obliterated). Both of the latter tend to involve only the anterior parts of the labium majus, but all these conditions extend to the groin. Whereas a hernia is as a rule reducible, a cyst of the canal of Nuck is not. Inguinal hernias usually disappear as pregnancy progresses, but varicoceles become worse. If a hernia contains bowel, it is resonant to percussion. A strangulated hernia will not be reducible, but the accompanying acute symptoms and the history should make the diagnosis clear.

*A haematoma* of the vulva may follow delivery or occur as the result of direct trauma. It is recognized as a bluish swelling, which is painful and tender and spreads up into the pelvis by the side of the vagina. The appearance is characteristic, and the diagnosis is made on the history.

An *endometrioma* is a rare cause of a blood-containing cyst on the vulva.
BENIGN NEW GROWTHS

- Caruncle
- Fibroma
- Fibromyoma
- Lipoma
- Hidradenoma
- Papilloma
- Lymphangioma
- Myxoma
- Angioma
- Melanoma
- Neuroma

As the vulva is comprised of skin, any swelling that can occur in a skin appendage can be found in the vulval region. Both fibromas and lipomas are seen in the vulva, and these may become pedunculated. They may occur at any age, are soft, oval or rounded, and are covered by vulval skin. They may grow slowly to reach the size of a fist. A lipoma is usually broader-based than a fibroma. Several other benign swellings are found on the vulva. They are usually solitary and small (about 1 cm or so in diameter), and their nature is arrived at by histology. A papilloma is a sessile benign tumour of the labile skin usually occurring in women of middle or old age. A hidradenoma is a tumour of sweat gland origin, which may be solid or cystic and can ulcerate. When ulcerated, it may clinically suggest the diagnosis of carcinoma; biopsy solves the problem. Less commonly are found fibromyoma, myxoma, angioma, lymphangioma, benign melanoma and neuroma, each distinguished by microscopic examination.

Tumours at the urethral meatus

_Urethral caruncles_ are frequent, especially in older women. A caruncle appears as a small, reddish, sessile growth arising from the posterior wall of the urethral meatus, causing bleeding and painful micturition. It is often very tender but may be symptomless. It is usually granulomatous, but it may be polypoidal and papillomatous. It has to be distinguished from prolapse of the urethral mucosa, in which there is a ring of protruding red tissue all round the urethral opening.

MALIGNANT NEW GROWTHS

- Squamous cell carcinoma
- Rodent ulcer
- Adenocarcinoma
- Sarcoma
- Melanoma
- Chorionic carcinoma

It must be emphasized that cancer within the vulva (Fig. V.16) is a very uncommon condition, and that any tumour that occurs in the skin can occur in the vulval region. The most common type is _squamous cell carcinoma_, which may have been preceded by pruritus but may be completely asymptomatic. It occurs mainly in postmenopausal women. Usually, it occurs as a single tumour, although on occasions it may present as kissing ulcers. The most common site is on the labia, and it spreads locally in the first instance and then to the inguinal lymph nodes. Squamous lesions account for 85 per cent of vulval cancers, the remainder comprising tumours of the skin and vulval appendages. Other malignant tumours found in the vulva include:

- Rodent ulcer (basal cell carcinoma), forming a flat plaque with its characteristic rolled edge
- Malignant melanoma (pigmented and non-pigmented)
- Adenocarcinoma arising in Bartholin's gland or in the urethra
- Sarcoma
- Undifferentiated tumours
- Metastatic tumours from primaries in the cervix, uterine body and ovary, which occur, though rarely
- Chorionic carcinoma

Figure V.16 Vulval carcinoma.
Ulceration can be defined as a persistent breach in any epithelial surface. The causes of ulceration include the following:

• Physical causes including pressure, chemicals (including urine), irradiation and scratching from intense itching. The vulval dystrophies may lead the woman to scratch sufficiently to breach the epithelium (see PRURITUS VULVAE, p. 529).

• Infective:
  – Sexually transmitted
  – Non-sexually transmitted

• Vascular insufficiency or compromise

• Sensory loss allowing ulceration due to trauma due to lack of sensation

• Malignancy produces a localized swelling that becomes ulcerated. The various types of malignancy are summarized under VULVA, SWELLING OF (p. 749). Premalignant lesions of the vulva do not usually cause ulceration of the vulva unless there has been scratching from pruritic symptoms.

INFECTIVE: SEXUALLY TRANSMITTED

Herpes
Primary infection occurs 2–7 days after inoculation with herpes simplex virus (HSV). Prodromal symptoms of tingling or itching are followed by vesicular eruptions, which rapidly erode, resulting in painful, shallow ulcers all over the vulva (Fig. V.17). They give rise to dysuria and, if secondarily infected, may lead to retention of urine, bilateral inguinal lymphadenopathy, fever and malaise. Herpes virus can be obtained from the vesicular fluid in the early stages, 85 per cent being due to the HSV type 2. The lesions, which are very painful, persist for 2–6 weeks before healing occurs and antibodies appear in the blood. The ulcers tend to recur at intervals of weeks or months, and the virus may be recovered from them. Coitus with a non-immune partner will pass on the infection. The disease is self-limiting in time, and the lesions eventually heal spontaneously. During pregnancy, the fetus is at risk if the episode is a primary infection.

Syphilis
Primary syphilis gives rise to an indurated ulcer that is characteristically painless unless it becomes secondarily infected. The incubation period is between 10 and 90 days following contact. Genital lesions in women often escape notice because they are hidden inside the vagina or on the cervix. The lesion has to be differentiated from an epithelioma. The serum from a chancre contains the spirochaete Treponema pallidum, which can be seen under a microscope with the aid of dark-ground illumination. If an epithelioma is suspected, the ulcer and swelling should be excised and examined microscopically. The chancre persists for 1–5 weeks, but serological tests for syphilis do not become positive for about 4–6 weeks after the appearance of the chancre. The serological tests most commonly performed are the Venereal Disease Research Laboratory (VDRL) slide test and the FTA-ABS (fluorescent treponemal antibody absorption) test, which have replaced the Wassermann, Kahn and TPI (treponemal immobilization) tests. To exclude primary syphilis, serological tests have to be done every week for 6 weeks after the appearance of the chancre. Between 2 weeks and 6 months after the chancre has healed, the generalized cutaneous eruption of secondary syphilis appears. Numerous moist, flat-topped papules occur on the vulva and round the anus. They are known as ‘condylomata lata’. In only one-third of untreated cases does tertiary syphilis occur, but not until some years after the primary lesion.

Lymphogranuloma venereum
This is a sexually transmitted disease found in the tropical and subtropical regions of Africa, Asia and south-eastern USA. It is due to a subtype of Chlamydia
trachomatis, and it begins with a vesicopustular eruption on the vulva, which soon disappears, but there follows painful suppuration in the inguinal nodes, with hypertrophy and ulceration of the groins, vulva and perineum. Later scarring may cause anal stricture or severe dyspareunia. The diagnosis may be made by isolating the organism, by the intradermal injection of virus antigen when a cutaneous reaction develops (Frei test), or by a complement fixation test.

**Granuloma inguinale**

This is a chronic venereal infective condition with a tendency to ulceration and massive granulation tissue affecting the vulva and groins. It is almost non-existent in Great Britain, but it is seen in India, Brazil and the West Indies, islands of the South Pacific, Australia, China and Africa. It starts as a raised papilloma, which soon ulcerates, the ulcer having a typical serpiginous outline. The granuloma in the groin rarely suppurates, but much scarring develops. Scrapings from the ulcers reveal the causative organism, the Donovan body, a small bacterium encapsulated in mononuclear leucocytes with a curved rod-like nucleus.

**Chancroid**

This is a very common cause of genital ulceration in tropical parts of the world, and occurs 3–5 days after coitus. It begins as a vesicopustule, which becomes a punched-out ulcer with a red base, or as a saucer-shaped, ragged ulcer. The lesion is extremely tender and produces a heavy foul discharge, which is contagious. It contains the causative organism, Ducrey's bacillus (*Haemophilus ducreyi*), a Gram-negative rod. The lesion may be solitary, or there may be several ulcers and associated painful inguinal adenitis, which may break down and discharge.

**Yaws**

This occurs in tropical countries and produces lesions similar in appearance to the condyloma lata of secondary syphilis.

**INFECTIVE: NON-SEXUALLY TRANSMITTED**

- **Aphthous ulcers**: these are analogous to the painful small ulcers that can be found in the mouth.
- **Tuberculosis**: this is a rare cause of vulval ulceration, but it may be associated with inguinal lymphadenopathy. The ulcers are highly indolent and can only be diagnosed with certainty on microscopic section.
- **Furunculosis**: these are boils caused by staphylococcal infection of the hair follicles. The condition is common and affects the labia majora in particular; shaving the area may predispose to this problem.
- **Diphtheria**: this condition produces ulceration with a characteristic membranous exudate.
- **Candidiasis**: mycotic and diabetic vulvitis due to *Candida* can cause soreness and pruritus of the vulva, with redness, excoriation and oedema of the skin, and a characteristic white curd-like discharge containing the mycelium of *Candida albicans*.

**SYSTEMIC DISEASE**

**Behçet's syndrome**

Behçet's syndrome is a rare autoimmune disorder of unknown cause that is characterized by oral and vulval ulceration. Iridocyclitis, arthritis and nervous system involvement are complications of severe cases. Diagnosis is difficult as there are no specific confirmatory tests (see www.behcets.com).

**Crohn's disease**

The vulva and perineum may be affected in up to 30 per cent of cases of Crohn's disease, and this may predate the gastrointestinal symptoms. The lesions appear like knife cuts in the skin. However, discharging sinuses and irregular ulcers are more common. This problem occurs very infrequently (see www.crohns.org.uk).

**Lipschutz ulcers**

These mainly occur on the labia minora and are of acute onset with an associated fever and lymphadenopathy. The cause is not known, but they may be due to Epstein–Barr virus.

**MALIGNANCY**

This has been discussed previously, but in summary the types of tumours that arise in this area are:

- Squamous carcinoma
- Melanoma
- Sarcoma
- Basal cell carcinoma
- Bartholin's glands adenocarcinoma
- Undifferentiated
- Possible secondary tumours

Biopsy will help in the tissue diagnosis, and an MRI scan may be useful to ascertain the extent of disease.
A weal is a transient erythematous lesion arising from the release of inflammatory mediators in the skin. It is a component of the Lewis triple response and may thus arise as a normal physiological response to firm stroking of the skin. In some susceptible individuals, there is an exaggeration of the normal wealing process, producing symptomatic dermographism, with wealing occurring from such innocent phenomena as the pressure of clothing on the skin. Some patients find this phenomenon alarming, but investigation is invariably unrewarding, and the response to oral antihistamines is somewhat variable.

Urticaria (Figs W.1 and W.2) is a common condition where weals arise in the skin spontaneously, presenting as randomly scattered, evanescent, erythematous lesions, sometimes with a rim of pallor at their margins, and sometimes associated with subcutaneous or mucosal swelling (angio-oedema) (Fig. W.3), which, if it affects the respiratory passages, may be life-threatening. Individual lesions may last for a matter of minutes or hours (normally not more than 24 hours), vanishing without trace, although new neighbouring weals may continue cropping. Weals can be produced experimentally by the intradermal injection of histamine, and they are largely blocked by antihistamines. The release of histamine from dermal mast cells is the suggested mechanism in most cases, although other inflammatory mediators may be involved (e.g. acetylcholine, kinins, prostaglandins and platelet activating factor).

Urticaria is very common (probably 10 per cent of the population will experience an attack at some stage) and, despite the clamour of sufferers for ‘allergy tests’, a careful history is the most important part of investigation. In acute urticaria, the onset is abrupt, and the condition may last from 1–2 days to several weeks. In most cases, there is a history of reaction to a food (e.g. strawberries, shellfish) or a drug. Peanut sensitivity can provoke especially violent episodes, often associated with angio-oedema or bronchospasm; in susceptible individuals, even a minute quantity may provoke an attack.

A very important form of urticaria, relatively recently recognized, is contact urticaria to latex. Rubber is produced from the sap of the tree Hevea brasiliensis, and prolonged exposure to rubber, especially rubber gloves in nurses or doctors, results in a type I immediate sensitivity. Itching and swelling start within minutes of contact and can be associated with severe systemic reactions. An allergen-specific IgE test to latex may support the diagnosis; latex-sensitive individuals must also avoid certain fruits – avocado, strawberry, kiwi fruit and banana – which contain cross-reactants. Chronic urticaria is usually idiopathic and may have an autoimmune basis. It may continue for years, but prophylactic antihistamines may help. Aspirin may
trigger flares, without itself being the primary cause. Antihypertensives of the angiotensin-converting enzyme inhibitor family may also be an unsuspected cause. If angio-oedema is a prominent feature, and there is a family history, the possibility of hereditary C₄ esterase inhibitor deficiency should be entertained. The diagnosis may be confirmed by a blood test; it is an important diagnosis, because attacks may be associated with abdominal pain and, if the true cause is known, a laparotomy may be avoided.

In some cases of urticaria, the triggering factor is a physical rather than an allergic one, and this is termed a physical urticaria (Fig. W.4). The condition is chronic, of unknown cause and often relatively resistant to treatment. An important example is cold-induced urticaria. Attacks may be triggered by, for example, eating an ice-cream; jumping into a cold swimming pool (or the sea) may be especially hazardous. The diagnosis may be confirmed by performing a provocation test with an ice cube. In delayed pressure urticaria, the attacks occur several hours after the event, and lesions arise in areas of the skin that have been subject to pressure (e.g. carrying a heavy shopping bag or walking on cobblestones). The episodes may last for some days and are associated with mild systemic upset. Other recognized triggers of physical urticaria include ultraviolet light, vibration and water; indeed, aquagenic urticaria is surprisingly common.

Cholinergic urticaria is a distinctive condition in which numerous intensely itchy pinhead papules develop some 10 minutes after sweating. Some patients can be induced to exhibit their symptoms by asking them to do mental arithmetic in the outpatient clinic. Susceptible individuals show an exaggerated reaction to the introduction of acetylcholine derivatives into the skin. Severe attacks are followed by a refractory period of 24–48 hours.

**SERUM SICKNESS**

Type III hypersensitivity causes weals of longer duration. Wealing is a rare feature of several general medical conditions and infections. In thyrotoxicosis and lymphoma, other signs and symptoms can be found but, in the early stages of systemic lupus erythematosus, urticaria may be the only clinical abnormality, the weals often persist for an unusually long time (2–4 days), and they may cause bruising. Viral hepatitis can begin with urticaria, the fever and icterus following 2–3 days later. A particular urticarial eruption appearing dramatically each evening often accompanies adult Still’s disease.

Weals can be the presenting feature of other dermatoses (e.g. pemphigoid, dermatitis herpetiformis or vasculitis). In a few atopic individuals, a contact urticaria occurs soon after the skin is touched by, for example, certain grasses, animal hair or saliva, or foods such as fish or fruits.

**WEIGHT GAIN**

Paul Carroll

The hypothalamus is central to the control of energy homeostasis in the body. This involves the control of hunger (via the lateral hypothalamic nucleus, LHA), which stimulates food intake, and satiety (via the ventromedial hypothalamic nucleus, VMH), which inhibits food intake. Body weight is influenced by the rate of energy expenditure, which is regulated by the secretion of hormones involved in the build-up of energy stores. Short-term signals for food intake affect the size and timing of individual meals. These involve internal sites in the gut and external stimuli such as food cues in the environment, as well as higher brain centres associated with the cognitive and emotional aspects of food intake. Long-term signals are mediated through the fat-derived hormone leptin, which gives a peripheral signal to the brain of adipose stores; leptin levels decrease in states of starvation. Neuropeptide Y (NPY) is present in the peripheral and central nervous systems, and it promotes anabolism by stimulating the secretion of insulin (independent of increased food intake), as well as stimulating adipose tissue lipoprotein lipase activity. Corticotrophin-releasing hormone has the opposite effect to NPY by having a catabolic effect.
The major causes of weight gain are highlighted in Box W.1.

Although it is usually simple to differentiate the major causes – especially where pregnancy or oedema is apparent – a scientific assessment of fat and muscle mass can be made by using skinfold thickness measurements, or by measuring electrical impedance. Overweight and obesity refer to conditions where body fat is in excess. The term ‘overweight’ is used when body weight is 110–119 per cent of the upper limit of an acceptable range (which is itself defined as 100 per cent), while the term ‘obesity’ refers to a body weight of 120 per cent or above. The acceptable range is based on life assurance figures of mortality, drawn up for an acceptable range (which is itself defined as 100 per cent of the upper limit of body weight) for the variations in energy expenditure reported.

A BMI of 25–29.9 is graded as overweight (Garrow Obesity Scale I), 30–40 as grade II obesity, and greater than 40 as morbidly obese (grade III).

Recent surveys have indicated the enormity of the problem in Western society. Both sexes show a rapid increase in weight in their mid-20s. Males tend to become progressively heavier until they reach their 50s, whereas in women, weight remains fairly static until the menopause, when there is a substantial weight gain. By the age of 25 years, about 30 per cent of males and females are overweight, whereas at 60 years of age about half have a weight problem considered to be a risk to health. In the 2001 census in the UK, 8 per cent of males and 12 per cent of females had a BMI of 30 or above. Although the risk to health increases with the degree of obesity, the configuration of adipose tissue may be an independent risk factor associated with an increased risk of hyperinsulinaemia, hyperlipidaemia, hypertension, ischaemic heart disease, cardiovascular events and death. A comparison of the circumference of waist and hip indicates that a ratio of over 0.8 in females and 0.95 in males is hazardous to health.

The cause of obesity in everyone is an energy intake in excess of energy expenditure. The scientific arguments arise over whether some individuals with a predisposition for obesity have a lower energy expenditure at the outset. Recent reports suggest a degree of variability in energy expenditure in any obese population and, if this is matched by a similar variability in appetite, this possibly explains in a simple manner the aetiology of obesity. Studies of twins indicate a degree of genetic involvement in the development of obesity, but there is no doubt that the cause is multifactorial, with environmental influences playing a major role. Much emphasis has been placed on an aetiological role for brown fat, which is a potent thermogenic tissue in rodents. Certainly, brown fat is present in adults, but it is thought to contribute little to daily energy expenditure and could not by itself account for the variations in energy expenditure reported.

Although the vast majority of obese individuals could be said to have ‘idiopathic’ or ‘simple’ obesity, it is always worthwhile considering the other rare causes of obesity (Box W.2).

Any hypothalamic lesion that destroys the ventromedial nucleus (classical ‘satiety’ area) may result in obesity. Other hypothalamic manifestations are often present in such patients. These include panhypopituitarism, Adiabetes insipidus, and also lethargy and somnolence. If the condition arises in childhood, the genitalia are usually poorly developed. In Fröhlich’s original case, a craniopharyngioma was pressing upon the hypothalamus, and hence such obesity associated with a hypogonadal state and poor growth is often referred to as Fröhlich’s syndrome. Most boys in whom the diagnosis of Fröhlich’s syndrome is entertained are in fact suffering from ‘simple’ obesity in which the pre-pubertal genitalia are buried in a pubic pad of fat.

Hypothyroidism leads to weight gain due both to a reduction in the metabolic rate and to the deposition of hydrophilic mucopolysaccharides all over the body, which causes fluid retention. Most cases are due to thyroid gland dysfunction and, in such patients, the serum thyroid-stimulating hormone (TSH) level is elevated. Hypothalamic pituitary lesions can result in a similar dysfunction (known as secondary hypothyroidism) with a low serum TSH.

<table>
<thead>
<tr>
<th>Box W.1 Causes of weight gain</th>
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<tbody>
<tr>
<td><strong>Excess fluid retention</strong></td>
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<tr>
<td>• Cardiac failure</td>
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<td>• Liver failure</td>
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<td>• Renal failure</td>
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<tr>
<td>• Nephrotic syndrome</td>
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<tr>
<td>• Periodic oedema</td>
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<tr>
<td>• Hypoproteinaemic states</td>
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<tr>
<td><strong>Lymphatic obstruction</strong></td>
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<td>• Milroy’s syndrome</td>
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<tr>
<td>• Elephantiasis</td>
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<tr>
<td>• Metastatic carcinoma</td>
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<tr>
<td><strong>Excess muscle</strong></td>
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<tr>
<td>• Precocious puberty</td>
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<tr>
<td>• Androgenic steroids</td>
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<tr>
<td>• Growth hormone</td>
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<tr>
<td>• Athletes, especially weightlifters</td>
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<tr>
<td><strong>Fat excess</strong></td>
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<tr>
<td>• Obesity</td>
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<tr>
<td><strong>Organ enlargement</strong></td>
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<tr>
<td>• Ovarian cyst</td>
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<td><strong>Pregnancy</strong></td>
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</table>
In most cases of thyrotoxicosis, weight decreases due to the rise in energy expenditure, hence resulting in an increase in weight.

**Pseudohypoparathyroidism** is associated with some degree of obesity. These patients are short, with stubby hands and feet due to the shortening of one or more (which can include all five) metacarpals and metatarsals. Ectopic calcification, including basal ganglion calcification, and cataracts have been reported. The cause appears to be due to a peripheral resistance to the metabolic effects of parathyroid hormone, producing serum hypocalcaemia. This is most likely to be a consequence of a deficiency in the membrane regulatory or coupling proteins (known as N or G). This membrane protein dysfunction can extend to other hormone receptors, hence explaining the abnormalities in secretion and in the effects of thyrotrophin, prolactin, gonadotrophin, vasopressin, glucagon and insulin sometimes encountered in this syndrome. There is also a variant disorder of similar phenotype, but normal biochemistry, termed **pseudo-pseudohypoparathyroidism**, which can occur independently or in the relatives of patients with pseudohypoparathyroidism.

**Cushing’s syndrome** is rare, but it can be differentiated from ‘simple’ obesity by its four cardinal signs: thin skin, conjunctival oedema (chemosis), frontal balding (most notably in women) and proximal myopathy. In a florid case, obesity is central, with thin arms and legs due to muscle wasting and a moon plethoric face (see Fig. F7). Oedema, hypertension, diabetes mellitus and severe osteoporosis are usually present. The latter often results in spontaneous rib and spine fractures, and in turn a loss of height and kyphosis. The latter condition must also be distinguished from the buffalo hump associated with fat over the upper thoracic spine. Acne, virilism, hirsutism and spontaneous bruising are often present. In children, this disease slows growth – unlike simple obesity, where growth is accelerated in the early years. Striae can be found in Cushing’s syndrome, but the absence does not exclude the condition. Striae are not pathognomonic of Cushing’s syndrome, as they are often found in healthy individuals who have had a rapid weight change.

In mild cases of Cushing’s syndrome, the differential diagnosis from simple obesity is not easy. If suspected, 24-hour urine samples should be measured for free cortisol, as this is elevated in Cushing’s syndrome. Some use an overnight 1 mg dexamethasone suppression test, the drug being taken at midnight and a blood cortisol value measured at 9 a.m. the next morning; a value of less than 100 nmol/l is considered normal. Although the

<table>
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<th>Box W2 Causes of obesity</th>
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<tr>
<td><strong>Idiopathic or simple obesity</strong></td>
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<tr>
<td>Energy intake in excess of energy expenditure; multifactorial causes</td>
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<tr>
<td><strong>Genetic</strong></td>
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<tr>
<td>Prader–Willi syndrome</td>
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<tr>
<td>Laurence–Moon–Biedl syndrome</td>
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<td>Alström’s syndrome</td>
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<td>Morel’s syndrome</td>
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<td>Morgagni syndrome</td>
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<td>Morgagni–Stewart–Morel syndrome</td>
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<td>Carpenter’s syndrome</td>
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<td>Cohen’s syndrome</td>
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<td>DIDMOAD</td>
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<tr>
<td><strong>Hypothalamic</strong></td>
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<tr>
<td>Trauma</td>
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<tr>
<td>Tumours</td>
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<tr>
<td>– Craniosphenoidya</td>
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<td>– Astrocytoma</td>
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<td>Inflammation</td>
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<td>– Meningitis</td>
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<td>– Encephalitis</td>
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<td>– Tuberculosis</td>
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<td>– Syphilis</td>
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<td>Infiltration</td>
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<td>– Sarcoiisis</td>
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<td>– Histiocytosis</td>
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<td><strong>Endocrine</strong></td>
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<td>Hypothalamus</td>
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<td>– Hypogonadotrophic hypogonadism</td>
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<td>– Growth hormone failure</td>
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<tr>
<td>Pituitary</td>
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<td>– Laron dwarfism</td>
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<td>– Hyperprolactinaemia</td>
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<td>– Cushing’s disease</td>
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<td>– Nelson’s syndrome</td>
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<tr>
<td>– Hypopituitarism (Fröhlich)</td>
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<td><strong>Thyroid</strong></td>
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<tr>
<td>Cretinism</td>
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<tr>
<td>Primary and secondary hypothyroidism</td>
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<tr>
<td>Rare cases of thyrotoxicosis</td>
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<tr>
<td><strong>Parathyroid</strong></td>
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<td>Pseudohypoparathyroidism</td>
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<tr>
<td>Pseudo-pseudohypoparathyroidism</td>
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<td><strong>Adrenal</strong></td>
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<tr>
<td>Cushing’s syndrome (ectopic source, adenoma, carcinoma)</td>
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<tr>
<td>Nesidioblastoma</td>
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<td>Insulinoma</td>
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<td>Beckwith–Wiedemann syndrome</td>
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<td><strong>Ovaries</strong></td>
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<td>Polycystic ovary syndrome</td>
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<td>Postmenopausal</td>
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<td><strong>Testes</strong></td>
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<tr>
<td>Primary hypogonadism</td>
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<td><strong>Inactivity</strong></td>
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<td>Mental retardation (e.g. Down’s syndrome)</td>
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<td>Physical disability (e.g. spina bifida)</td>
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<td><strong>Drugs</strong></td>
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<td>Insulin</td>
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<td>Cessation of smoking</td>
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<td>Sulphonylurea agents</td>
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<td>Corticosteroids</td>
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<td>Oestrogen</td>
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<tr>
<td>Alcohol excess (pseudo-Cushing’s syndrome)</td>
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<tr>
<td>Cyproheptadine</td>
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<tr>
<td>Abnormal fat distribution</td>
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<tr>
<td>Multiple lipomatosis</td>
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<tr>
<td>Partial lipodystrophy</td>
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<tr>
<td>Painful fat</td>
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<td>Dercum’s disease</td>
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and an absent or delayed TSH response to thyroid-releasing hormone. Weight loss can be rapid following the introduction of replacement thyroxine therapy in severely hypothyroid patients. Nevertheless, in those whose obesity precedes the development of mild hypothyroidism, weight loss can be slight with thyroid replacement. This is a source of disappointment to many, but indicative of an underlying ‘simple’ obese problem. In most cases of thyrotoxicosis, weight decreases due to an increase in energy expenditure. In some rare cases of thyrotoxicosis, the increase in appetite produces an energy intake in excess of the rise in energy expenditure, hence resulting in an increase in weight.
WEIGHT GAIN

simple overnight dexamethasone test is helpful, no test is ever totally reliable and, if clinical suspicion persists, further investigations should be carried out on an inpatient basis. Alcohol can mimic Cushing’s syndrome both in clinical features and by increasing urine free cortisol output; this situation is termed pseudo-Cushing’s. Admission to hospital with a total ban on alcohol will eventually result in normal cortisol values.

Patients with insulinoma are said to be mildly obese as insulin stimulates appetite. Hyperinsulinaemia due to islet cell hyperplasia can occur in the rare abnormality of Wiedemann–Beckwith syndrome. These children grow more rapidly, have an enlarged tongue, and omphalocele, and tend to be mildly obese.

In polycystic ovarian syndrome, over half the patients are obese. The reason for this is not known, but obesity tends to perpetuate the syndrome because of the conversion in adipose tissue of ovarian androstenedione to oestrone. The other features of the syndrome are oligomenorrhea or menorrhagia, infertility, hirsutism and enlarged, polycystic ovaries. Biochemically, serum luteinizing hormone is elevated in comparison to follicle-stimulating hormone, associated with a raised serum testosterone and androstenedione. Many have wondered whether hyperplactinaemia induces weight gain, as the clinical impression is that many such women are obese. One survey disputed whether obesity was more prevalent in hyperprolactinaemia, but the impression still remains.

Males with hypogonadism of whatever cause tend to be slightly obese. The distribution of fat is of a female pattern over the lower abdomen, hips and thighs. Growth hormone failure in children is associated with mild obesity, as is the rare Laron dwarf who has decreased levels of insulin-like growth factors and insulin-like growth factor-binding protein-3 (see STATURE, SHORT, p. 639).

There are various other congenital syndromes associated with obesity. Prader–Willi syndrome is characterized by muscular hypotonia, short stature, small body and feet, mental retardation, hypogonadism and gross obesity. Hyperappetite appears to be the principal cause of their obesity, although a hypothalamic thermic abnormality has been reported. Recently, deletions of chromosome 15 have been implicated as the cause of this condition. In Laurence–Moon–Biedl syndrome, obesity is associated with short stature (in some), retarded sexual development, mental retardation, retinitis pigmentosa, and either polydactyly or syndactyly.

In Alstrom’s syndrome, there is obesity, diabetes mellitus, nerve deafness, retinal degeneration and cataracts, producing childhood blindness and late-onset nephropathy. Many patients with Down’s syndrome are obese, as are many children with mental or physical retardation, possibly due to inactivity associated with too high an energy intake.

Morgagni–Stewart–Morel syndrome is a combination of the syndromes described by Morel and Morgagni. In Morel’s syndrome, there is obesity, hyperostosis of the frontal bone and headache. In Morgagni syndrome, the obesity is associated with internal frontal hyperostosis and virilism.

Carpenter’s syndrome is associated with acrocephaly, polydactyly and syndactyly, mental retardation, male hypogonadism and mild obesity. In Cohen’s syndrome, obesity is associated with severe mental retardation, microcephaly and short stature and facial abnormalities.

DIDMOAD is an acronym for the major features of the syndrome consisting of diabetes insipidus, diabetes mellitus, optic atrophy and deafness. In most such patients, this is associated with mild obesity, as well as bladder and ureter atonia.

EFFECTS OF DRUG INTAKE ON OBESITY

Some drugs may cause obesity or a worsening of a pre-existing weight problem. Insulin given to those with diabetes tends to lead to weight gain, as do the sulphonylurea drugs, but usually this is not seen with the biguanide metformin. Weight gain noted when glycaemic control is improved with insulin is partly due to a reduction in both energy expenditure and urine glucose losses. Glucocorticoids in excess rapidly cause weight gain and produce iatrogenic Cushing’s syndrome. Oestrogen can also induce weight gain. Although water retention has been implicated, a reduction in energy expenditure in the second half of the menstrual cycle due to a lack of ovulation also plays a role. A frequent finding is of weight gain on cessation of smoking. Recent studies have shown that a smoked cigarette increases sympathetic drive, and on average one cigarette increases energy expenditure by about 9 kcal. Hence, if smoking is ceased, energy intake must be appropriately reduced to prevent weight gain. Nevertheless, many people tend to overeat on cessation of smoking, which exacerbates the problem.

BODY FAT DISTRIBUTION

An abnormal distribution of fat occurs in lipodystrophies; these comprise a group of disorders ranging from generalized lipoatrophy to partial forms. Partial lipoatrophy (lipodystrophia progressiva) is characterized by a symmetrical loss of subcutaneous fat over dermatome areas of the face and upper body, with a normal or even excessive amount of adipose tissue.
below the waist. Occasionally, the atrophy is confined below the waist without upper body involvement. This condition, which mainly affects females, is associated with hyperglycaemia, hyperinsulinaemia, marked insulin resistance, hyperlipaemia and hepatosplenomegaly. In Dercum's disease, obesity is generalized, but there is pain and tenderness in the more prominent fatty deposits, for reasons as yet unknown.

Abnormal body fat distribution may occur in patients with AIDS. Dorsocervical fat pad enlargement (buffalohump), benign symmetrical lipomatosis, abdominal girth enlargement and breast hypertrophy (see WEIGHT LOSS, ‘HIV infection’, p. 760) have all been documented in patients with AIDS. It remains unclear whether this change in body fat distribution is related to or aggravated by drug therapy (e.g. with the protease inhibitors), as some aspects of the change in body fat distribution have been documented in patients not receiving therapy. The redistribution of body fat occurs in association with peripheral and gluteal wasting; although this is a very similar clinical picture to that seen in Cushing's syndrome, the cortisol biochemistry is not abnormal.

**WEIGHT LOSS**

Mark Kinirons & Duncan Churchill

Loss of weight may be the result of inadequate food intake, Absorption or retention. It may also be due to increased utilization. It is important to distinguish between those patients who lose weight in spite of normal food intake and those whose calorie intake is diminished. In children, the most common causes are malnutrition from injudicious feeding, gastrointestinal disease and infections (see MARASMUS, p. 398). In adults, when loss of weight is considerable, one thinks first of malignant disease, infection such as HIV, *tuberculosis*, or endocrine abnormalities such as hyperthyroidism and diabetes mellitus. Without any evidence of organic disease, other conditions to consider include depression and an eating disorder.

Research indicates that, in 35 per cent of patients losing weight, no specific cause is usually found. In the remainder, a serious underlying cause is discovered, with 10 per cent having a psychiatric aetiology, the others having a physical disorder, the most common being cancer, gastrointestinal disease, heart failure, alcohol abuse, obstructive airways disease, poorly controlled diabetes mellitus, thyrotoxicosis and HIV. Taking a good history and performing a thorough physical examination will provide direction on the area for investigation. The single most helpful investigation is a chest radiograph, which often reveals masses, infiltrates (e.g. opportunistic infection), heart failure or lymph node enlargement.

**WEIGHT LOSS WITH ADEQUATE FOOD INTAKE**

1. Increased utilization
   - Hyperthyroidism
   - Chronic infections (e.g. *pulmonary tuberculosis*)
   - Anxiety states, food phobias
   - Drugs: L-thyroxine, amphetamine

2. Diminished absorption
   - Intestinal hypermotility states
   - Chronic pancreatitis
   - Carcinoid
   - Gluten enteropathy
   - Short-circuit operations
   - Post-colectomy and post-gastrectomy states
   - Chronic hepatic disease
   - Dysphagia (e.g. *scleroderma*)
   - Whipple's disease
   - Lymphatic obstruction
   - Drugs: serotonin reuptake inhibitors (paroxetine, fluoxetine, citalopram); sibutramine and orlistat, both used in the treatment of obesity

3. Abnormal calorie loss
   - Diabetes mellitus
   - Fistulas
   - Intestinal parasites

**WEIGHT LOSS WITH DIMINISHED FOOD INTAKE**

1. Psychogenic
   - Depression
   - Anorexia nervosa
   - Psychoses

2. Gastrointestinal
   - Gastric ulcer
   - Malignancy
   - Chronic colitis
   - Hepatobiliary disease

3. Malignant conditions
   - Lymphoma
   - Leukaemia
   - Carcinoma: adenocarcinoma/squamous-cell carcinoma
   - Sarcoma

4. Uraemia

5. Chronic infections

6. Chronic non-infective inflammatory conditions
   - Rheumatoid arthritis
   - Systemic lupus erythematosus
   - Dermatomyositis
   - Polymyositis nodosa
   - Giant-cell arthritis
   - Systemic sclerosis
7. Chronic intoxications
   - Alcohol
   - Addictive drugs
   - Heavy smoking
   - Lead
8. Endocrine disease
   - Addison's disease and some cases of hypopituitarism (Simmond's disease)
   - Phaeochromocytoma
   - Gut hormone tumours (e.g. vipoma)
9. Food intolerance
10. Chronic cardiac conditions
11. Chronic lung conditions (e.g. obstructive airways disease)
12. HIV infection

Any condition of the gastrointestinal tract that interferes with the intake, digestion and absorption of food may produce loss of weight; for example, gastrointestinal neoplasia anywhere from the oral cavity to the anal canal; gastritis or gastric ulcer, inflammatory bowel disease and bowel surgery; and partial or total gastrectomy, small bowel surgery or colectomy. 

Chronic infections may produce loss of weight by interfering with general nutrition. This is seen in many who have returned from the tropics after infection by dysentery, yellow fever, malaria, dengue or hepatitis. The chronic infection of a joint, the skin or the renal tract may produce loss of weight in a similar way. 

Liver disorders may affect general nutrition, and loss of weight may occur in some sufferers from cirrhosis; loss of body weight may be masked by the weight gain due to ascites. Malignant change in a cirrhotic liver is not uncommon. Hepatocellular carcinoma may develop in patients with haemochromatosis and hepatitis C virus infection. In drug addiction and HIV (see below), weight loss is often considerable. 

The effect of alcohol upon body weight is variable, with some persons becoming stout, others thin, and others changing little. This depends greatly on food (calorie) intake, but beer-drinkers tend to become obese and pot-bellied. Broadly speaking, it is heavy spirit drinkers who lose weight, and in some cases serious doubts may arise whether the weight loss in such a patient is due to alcoholic habits alone or whether there is underlying neoplasia or tuberculous infection. When alcoholism leads to peripheral neuropathy, there is often a rapid and extreme loss of weight. In chronic debilitating disorders such as rheumatoid arthritis, multiple sclerosis or hemiplegia, a marked loss of weight may occur, as it may in chronic congestive heart failure, where oedema may mask the wasting.

Figure W.5 Crease pigmentation indicated Addison's disease in this patient, who had noted progressive and rapid weight loss associated with diarrhoea and lassitude.

The loss of weight in old age, due to diminished intake of and lessened interest in food, is usually gradual and slow: if otherwise, neoplasia or chronic infection, such as tuberculosis, depression, giant-cell arteritis or some other cause should be suspected. 

Diabetes mellitus, especially in the young, may have loss of weight as its earliest symptom. In Addison's disease, the loss of weight may be marked. There may or may not have been attacks of syncope or of diarrhoea. The diagnosis is suggested by brown pigmentation of the skin, particularly in the flexures and groins (Fig. W.5), and also beneath the mucous membranes – particularly of the mouth, inside the lips or within the cheeks – where it is a grey colour. The blood pressure is usually low, and shows a marked postural fall.

Loss of weight is a prominent feature in cases of hyperthyroidism; indeed, it may be the first symptom to attract attention, preceding those of tachycardia, nervousness, excessive perspiration, fine tremor of the outstretched fingers, exophthalmos and symmetrical enlargement of the thyroid gland. Anorexia nervosa is a condition in which wasting is the prominent symptom, but amenorrhoea often occurs very early in the disease.

HIV INFECTION

Between 11 and 18 per cent of AIDS patients suffer from the wasting syndrome, which is an involuntary loss of 10 per cent or more of their body weight. The wasting that occurs in AIDS results from a combination of abnormalities, including hypermetabolism, infection, malabsorption and anorexia. It remains unclear whether the testosterone deficiency found in patients with AIDS contributes to the weight loss and
muscle wasting. The wasting syndrome is associated with progressive HIV infection, and contributes directly to mortality. During the evaluation of weight loss in patients with AIDS, mucosal, gastrointestinal and systemic infection, as well as malignancy and hypogonadism, must be searched for. A rapid loss of weight is usually associated with active secondary infection. The compensatory decrease in resting energy expenditure (REE) that usually occurs during decreased caloric intake does not take place in AIDS patients, who appear to maintain a high REE in the presence of a decreased calorie intake. A syndrome of lipodystrophy characterized by subcutaneous fat loss in the face and extremities has been observed in people with HIV treated with antiretroviral drugs, particularly the nucleoside analogue stavudine.

Polyphonic wheeze present during tidal breathing is a reliable sign of severe airways obstruction. Normal subjects can generate polyphonic wheezes, but only on forced expiratory effort.

**FIXED MONOPHONIC WHEEZE**
When a bronchus is narrowed by stenosis of an intrabronchial tumour, a low-pitched monophonic wheeze may be heard, often on inspiration, in association with noisy breathing. The low note of the wheeze is related to the tumour mass, which is set in slow oscillation by high velocity gas flow. The pitch can be varied within a narrow range by altering the velocity of gas flow. Stridor is a special example of this sound.

**RANDOM MONOPHONIC WHEEZE**
A particular variety of wheeze distinct from the polyphonic expiratory sounds may be heard in widespread airflow obstruction, overlapping throughout inspiration and expiration, and with varying duration, timing and pitch.

**SEQUENTIAL INSPIRATORY WHEEZE**
In patients with pulmonary oedema or inflammatory infiltration, a brief high-pitched wheeze can frequently be heard late in inspiration in association with inspiratory crackles. The musical note may repeat from breath to breath, or disappear and reappear at different times. These sounds occur in deflated areas of lung and are classically heard in extrinsic allergic alveolitis, also known as hypersensitivity pneumonitis. They have been described as ‘squeaks and squarks’.

The paradoxical absence of wheezing may be of great clinical importance, indicating severe and widespread airflow obstruction. The production of a wheeze requires an airway on the point of closure and an optimum velocity of gas flow at the site of stenosis to set the bronchial walls in oscillation. In patients with severe ventilatory failure, wheezing may be absent because the velocity of flow is too slow to oscillate the airways on the point of closure. For the same reasons, deteriorating asthmatic patients may become less wheezy and eventually develop a silent chest, indicating ominous airways obstruction.

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**WHEEZE**
Alex West
(See also STRIDOR, p. 652.)

Wheezes are continuous, musical sounds, typically loudest during expiration, that may be heard at the mouth or with the aid of a stethoscope. These sounds are generated by vibration of an airway, not only narrowed but almost closed to allow the walls to touch lightly. Air accelerating through the narrowed airway generates pressure fluctuations causing the airway walls to oscillate rapidly, producing a musical wheeze. The pitch of the wheeze depends upon the speed of vibration rather than the length or calibre of the airway. Four clinical types of wheeze may be identified: expiratory polyphonic, fixed monophonic, random monophonic and sequential inspiratory.

**EXPIRATORY POLYPHONIC WHEEZE**
This is a complex musical sound commonly associated with widespread airflow obstruction due to chronic obstructive pulmonary disease or bronchial asthma. The wheeze, together with a background of loud, noisy breathing, is audible at the mouth. When listened to on the chest wall, the higher frequencies of the breath sounds are filtered out, and the wheezes dominate.
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For nearly a century, *French’s Index of Differential Diagnosis* has been invaluable in assisting clinicians to quickly and correctly diagnose a disease from a whole range of presenting symptoms. Arranged alphabetically by presenting symptom, the text helps readers to identify each presentation, describes the different diagnoses that it could represent, and explains the signs and tests used to make a diagnosis.

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The 16th Edition of this essential classic text offers a succinct and well-illustrated aide-memoire; it remains indispensable to trainee and established doctors in both general and hospital practice and an invaluable reference for medical students.

**MARK T. KINIRONS** BSc Hons MD FRCPI FRCP Consultant Physician and Honorary Senior Lecturer, Department of Ageing and Health, Guy’s and St Thomas’ Hospitals, London, UK

**HAROLD ELLIS** CBE DM MCh FRCS Emeritus Professor of Surgery, Division of Anatomy, Cell and Human Biology, Guy’s, King’s and St Thomas’ School of Biomedical Sciences, London, UK