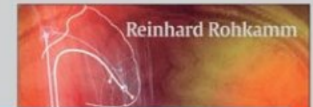
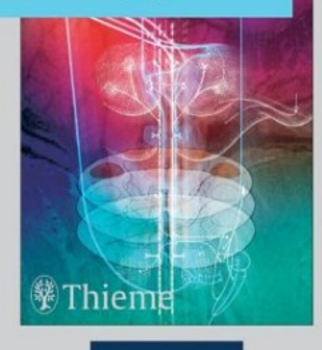
THIEME FLEXIBOOK



Color Atlas of Neurology





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Preface

The nervous system and the muscles are the seat of many primary diseases and are affected secondarily by many others.

This pocket atlas is intended as an aid to the detection and diagnosis of the symptoms and signs of neurological disease. The text and illustrations are printed on facing pages, to facilitate learning of the points presented in each.

The book begins with a summary of the fundamentals of neuroanatomy in Chapter 1. Chapter 2 concerns the functions of the nervous system and the commonly encountered syndromes in clinical neurology. Individual neurological diseases are discussed in Chapter 3. The clinical neurological examination is best understood once the material of the first three chapters is mastered; it is therefore presented in the last chapter, Chapter 4.

The choice of topics for discussion is directed toward questions that frequently arise in clinical practice. Some of the illustrations have been reproduced from previous works by other authors, because they seemed to us to be optimal solutions to the problem of visually depicting a difficult subject. In particular, we would like to pay tribute here to the graphic originality of the late Dr. Frank H. Netter.

Many people have lent us a hand in the creation of this book. Our colleagues at the Sanderbusch Neurological Clinic were always ready to help us face the difficult task of getting the book written while meeting the constant demands of patient care. I (R.R.) would particularly like to thank our Oberärzte (Senior Registrars), Drs. Helga Best and Robert Schumann, for their skillful cooperation and support over several years of work. Thanks are also due to the radiologists, Drs. Benno Wördehoff and Ditmar Schönfeld, for providing images to be used in the illustrations. This book would never have come about without the fascination for neurology that was instilled in me in all the stages of my clinical training; I look back with special fondness on the time I spent as a Resident in the Department of Neurology at the University of New Mexico (Albuquerque). Above all, I thank the many patients, past and present, who have entrusted me with their care.

Finally, cordial thanks are due to the publishers, Georg Thieme Verlag, for their benevolent and surefooted assistance throughout the development of this book, and for the outstanding quality of its production. Among the many members of the staff to whom we are grateful, we would like to single out Dr. Thomas Scherb, with whom we were able to develop our initial ideas about the format of the book, as well as Dr. Clifford Bergman and Gabriele Kuhn, who saw this edition through to production with assurance, expertise, and the necessary dose of humor.

We dedicate this book to our families: Christina, Claire, and Ben (R.R.) and Birgit, Jonas, and Lukas (M.G.).

Reinhard Rohkamm, Sande Manfred Güther, Bermatingen Autumn 2003

Contents

1 Fundamentals

Overview	2
Skull	4
Meninges	6
Cerebrospinal Fluid	8
Blood Vessels	10
Carotid Arteries	11
Anterior Circulation of the Brain	12
Vertebral and Basilar Arteries	14
Posterior Circulation of the Brain	16
Intracranial Veins	18
Extracranial Veins	20
Spinal Circulation	22

Central Nervous Sysstem	24
Anatomical and Functional Organization .	25
Brain Stem	26
Cranial Nerves	28
Spine and Spinal Cord	30
Peripheral Nervous System	32
Dermatomes and Myotomes	33
Brachial Plexus	34
Nerves of the Upper Limb	35
Lumbar Plexus	36
Nerves of the Lower Limb	37

2 Normal and Abnormal Function of the Nervous System 39

Motor Function	40
Reflexes	41
Motor Control	42
Motor Execution	44
Central Paralysis	46
Peripheral Paralysis	50
Cerebellum	54
Vestibular System	56
Vertigo	58
Gait Disturbances	60
Tremor	62
Dystonia	64
Chorea, Ballism, Dyskinesia, Myoclonus	66
Myoclonus, Tics	68
Brain Stem Syndromes	70
Midbrain Syndromes	71
Pontine Syndromes	72
Medullary Syndromes	73

74
75
76
78
80
82
84
86
88
90
92
94
96
98
00
02
04
06

Contents

Pain	108
Sleep Normal Sleep Sleep Disorders	113

Disturbances of Consciousness	116
Acute Disturbances of Consciousness	116
Coma	118
Comalike Syndromes, Death	120

Behavioral Manifestations of Neurological

Disease	122
Language	124
Aphasia	126
Agraphia, Alexia, Acalculia, Apraxia	128
Speech Disorders	130

Disturbances of Orientation	132
Disturbances of Memory	134
Dementia	136
Pseudo-neurological Disorders	138
Autonomic Nervous System (ANS)	140
Organization	141
Hypothalamus	142
Limbic System and Peripheral ANS	144
Heart and Circulation	148
Respiration	150
Thermoregulation	152
Gastrointestinal Function	154
Bladder Function, Sexual Function	156
Intracranial Pressure	158

3 Neurological Syndromes

Central Nervous System	166
Stroke	167
Headache	182
Epilepsy: Seizure Types	192
Epilepsy: Classification	196
Epilepsy: Pathogenesis and Treatment	198
Nonepileptic Seizures	200
Parkinson Disease: Clinical Features	206
Parkinson Disease: Pathogenesis	210
Parkinson Disease: Treatment	212
Multiple Sclerosis	214
CNS Infections	222
Brain Tumors	254

266
276
282
288
296
306
316
316

4 Diagnostic Evaluation

Diagnostic Evaluation	350
History and Physical Examination	350
Neurophysiological and Neuro-	
psychological Tests	352

Cerebrovascular Ultrasonography,	
Diagnostic Imaging, and Biopsy	
Procedures	353

5 Appendix

References	09	Index	415

165

349

355

Fundamentals

- Anatomy
- Physiology

Neurology is the branch of medicine dealing with diseases of the central, peripheral, and autonomic nervous systems, including the skeletal musculature.

Central Nervous System (CNS)

Brain

The **forebrain** or prosencephalon (supratentorial portion of the brain) comprises the telencephalon (the two cerebral hemispheres and the midline structures connecting them) and the diencephalon.

The **midbrain** or mesencephalon lies between the fore brain and the hind brain. It passes through the tentorium cerebelli.

The **hindbrain** or rhombencephalon (infratentorial portion of the brain) comprises the pons, the medulla oblongata (almost always called "medulla" for short), and the cerebellum. The mid brain, pons, and medulla together make up the **brain stem**.

Spinal cord

The spinal cord is approximately 45 cm long in adults. Its upper end is continuous with the medulla: the transition is defined to occur just above the level of exit of the first pair of cervical nerves. Its tapering lower end, the conus medullaris, terminates at the level of the L3 vertebra in neonates, and at the level of the L1-2 intervertebral disk in adults. Thus, lumbar puncture should always be performed at or below L3-4. The conus medullaris is continuous at its lower end with the threadlike filum terminale, composed mainly of glial and connective tissue, which, in turn, runs through the lumbar sac amidst the dorsal and ventral roots of the spinal nerves, collectively called the cauda equina ("horse's tail"), and then attaches to the dorsal surface of the coccyx. The cervical, thoracic, lumbar, and sacral portions of the spinal cord are defined according to the segmental division of the vertebral column and spinal nerves.

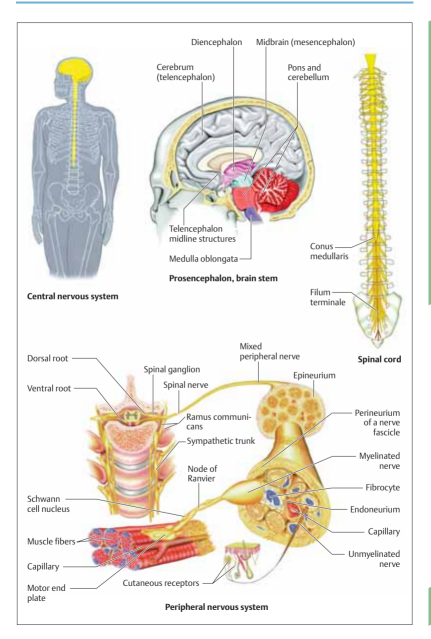
Peripheral Nervous System (PNS)

The peripheral nervous system connects the central nervous system with the rest of the body. All motor, sensory and autonomic nerve cells and fibers outside the CNS are generally considered part of the PNS. Specifically, the PNS comprises the ventral (motor) nerve roots, dorsal (sensory) nerve roots, spinal ganglia, and spinal and peripheral nerves, and their endings, as well as a major portion of the autonomic nervous system (sympathetic trunk). The first two cranial nerves (the olfactory and optic nerves) belong to the CNS, but the remainder belong to the PNS.

Peripheral nerves may be purely motor or sensory but are usually mixed, containing variable fractions of motor, sensory, and autonomic nerve fibers (axons). A peripheral nerve is made up of multiple bundles of axons, called fascicles, each of which is covered by a connective tissue sheath (perineurium). The connective tissue lying between axons within a fascicle is called endoneurium, and that between fascicles is called epineurium. Fascicles contain myelinated and unmvelinated axons, endoneurium, and capillaries. Individual axons are surrounded by supportive cells called Schwann cells. A single Schwann cell surrounds several axons of unmyelinated type. Tight winding of the Schwann cell membrane around the axon produces the myelin sheath that covers myelinated axons. The Schwann cells of a myelinated axon are spaced a small distance from one another; the intervals between them are called nodes of Ranvier. The nerve conduction velocity increases with the thickness of the myelin sheath. The specialized contact zone between a motor nerve fiber and the muscle it supplies is called the neuromuscular junction or motor end plate. Impulses arising in the sensory receptors of the skin, fascia, muscles, joints, internal organs, and other parts of the body travel centrally through the sensory (afferent) nerve fibers. These fibers have their cell bodies in the dorsal root ganglia (pseudounipolar cells) and reach the spinal cord by way of the dorsal roots.

Autonomic Nervous System (ANS)

The autonomic nervous system regulates the function of the internal organs in response to the changing internal and external environment. It contains both central (p. 140 ff) and peripheral portions (p. 146ff).



The skull (cranium) determines the shape of the head; it is easily palpated through the thin layers of muscle and connective tissue that cover it. It is of variable thickness, being thickre and sturdier in areas of greater mechanical stress. The thinner bone in temporal and orbital portions of the cranium provides the so-called *bone windows* through which the basal cerebral arteries can be examined by ultrasound. Thinner portions of the skull are more vulnerable to traumatic fracture. The only joints in the skull are those between the auditory ossicles and the *temporomandibular joints* linking the skull to the jaw.

Neurocranium

The neurocranium encloses the brain, labyrinth, and middle ear. The outer and inner tables of the skull are connected by cancellous bone and marrow spaces (*diploë*). The bones of the *roof of the cranium* (calvaria) of adolescents and adults are rigidly connected by sutures and cartilage (synchondroses). The *coronal suture* extends across the frontal third of the cranial roof. The *sagittal suture* lies in the midline, extending backward from the coronal suture and bifurcating over the occiput to form the *lambdoid suture*. The area of junction of the frontal, parietal, temporal, and sphenoid bones is called the *pterion*; below the pterion lies the bifurcation of the middle meningeal artery.

The inner skull base forms the floor of the cranial cavity, which is divided into anterior, middle, and posterior cranial fossae. The *anterior fossa* lodges the olfactory tracts and the basal surface of the frontal lobes; the *middle fossa*, the basal surface of the temporal lobes, hypothalamus, and pituitary gland; the *posterior fossa*, the cerebellum, pons, and medulla. The anterior and middle fossae are demarcated from each other laterally by the posterior edge of the (lesser) wing of the sphenoid bone, and medially by the *jugum sphenoidale*. The middle and posterior fossae are demarcated from each other laterally by the upper rim of the petrous pyramid, and medially by the dorsum sellae.

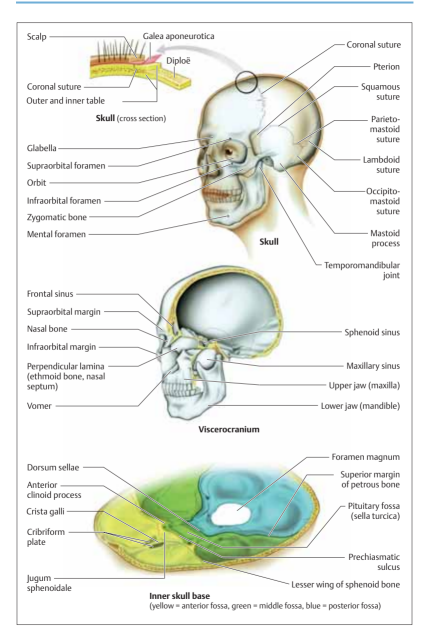
Scalp

The layers of the scalp are the skin (including epidermis, dermis, and hair), the subcuticular connective tissue, the fascial galea aponeurotica, subaponeurotic loose connective tissue, and the cranial periosteum (pericranium). The hair of the scalp grows approximately 1 cm per month. The connection between the galea and the pericranium is mobile except at the upper rim of the orbits, the zygomatic arches, and the external occipital protuberance. Scalp injuries superficial to the galea do not cause large hematomas, and the skin edges usually remain approximated. Wounds involving the galea may gape; scalping injuries are those in which the galea is torn away from the periosteum. Subgaleal hemorrhages spread over the surface of the skull.

Viscerocranium

The viscerocranium comprises the bones of the orbit, nose, and paranasal sinuses. The superior margin of the orbit is formed by the frontal bone, its inferior margin by the maxilla and zygomatic bone. The frontal sinus lies superior to the roof of the orbit, the maxillary sinus inferior to its floor. The nasal cavity extends from the anterior openings of the nose (nostrils) to its posterior openings (choanae) and communicates with the paranasal sinuses-maxillary, frontal, sphenoid, and ethmoid. The infraorbital canal, which transmits the infraorbital vessels and nerve, is located in the superior (orbital) wall of the maxillary sinus. The portion of the sphenoid bone covering the sphenoid sinus forms, on its outer surface, the bony margins of the optic canals, prechiasmatic sulci, and pituitary fossa.

Skull



Meninges

The *meninges* lie immediately deep to the inner surface of the skull and constitute the membranous covering of the brain. The pericranium of the inner surface of the skull and the dura mater are collectively termed the *pachymeninges*, while the pia mater and arachnoid membrane are the *leptomeninges*.

Pachymeninges

The pericranium contains the meningeal arteries, which supply both the dura mater and the bone marrow of the cranial vault. The pericranium is fused to the dura mater, except where they separate to form the dural venous sinuses. The virtual space between the pericranium and the dura mater-the epidural space-may be forced apart by a pathological process, such as an epidural hematoma. Immediately beneath the dura mater, but not fused to it, is the arachnoid membrane; the intervening virtual space-the subdural space-contains capillaries and transmits bridging veins, which, if injured, can give rise to a subdural hematoma. The falx cerebri separates the two cerebral hemispheres and is bordered above and below by the superior and inferior sagittal sinuses. It attaches anteriorly to the crista galli, and bifurcates posteriorly to form the tentorium cerebelli, with the straight sinus occupying the space between the falx and the two halves of the tentorium. The much smaller falx cerebelli separates the two cerebellar hemispheres; it encloses the occipital sinus and is attached posteriorly to the occipital bone.

The tentorium cerebelli separates the superior aspect of the cerebellum from the inferior aspect of the occipital lobe. It rises toward the midline, taking the shape of a tent. The opening between the two halves of the tentorium, known as the tentorial *notch* or *incisura*, is traversed by the midbrain; the medial edge of the tentorium is adjacent to the midbrain on either side. The tentorium attaches posteriorly to the sulcus of the transverse sinus, laterally to the superior rim of the pyramid of the temporal bone, and anteriorly to the anterior and posterior clinoid processes. The tentorium divides the cranial cavity into the *supratentorial* and *infratentorial spaces*.

The pituitary stalk, or infundibulum, accompanied by its enveloping arachnoid membrane, passes through an aperture in the posterior portion of the *diaphragma sellae* (diaphragm of the sella turcica), a horizontal sheet of dura mater lying between the anterior and posterior clinoid processes. The pituitary gland itself sits in the sella turcica, below the diaphragm.

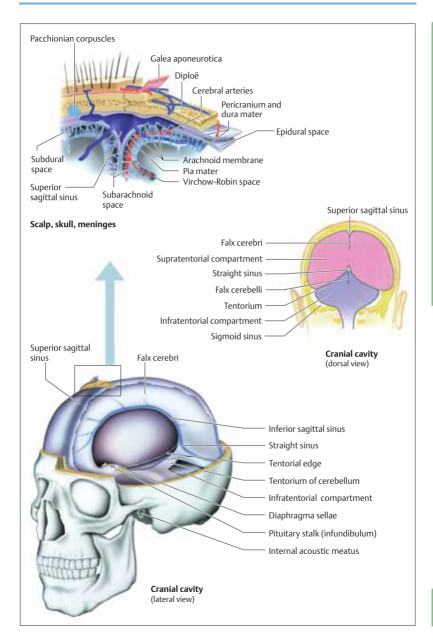
The meningeal branches of the three divisions of the trigeminal nerve (pp. 28 and 94) provide sensory innervation to the dura mater of the cranial roof, anterior cranial fossa, and middle cranial fossa. The meningeal branch of the vagus nerve (p. 29), which arises from its superior ganglion, provides sensory innervation to the dura mater of the posterior fossa. Pain can thus be felt in response to noxious stimulation of the dura mater, while the cerebral parenchyma is insensitive. Some of the cranial nerves, and some of the blood vessels that supply the brain, traverse the dura at a distance from their entry into the skull, and thereby possess an intracranial extradural segment, of a characteristic length for each structure. Thus the rootlets of the trigeminal nerve, for instance, can be approached surgically without incising the dura mater.

Pia Mater

The cranial pia mater is closely apposed to the brain surface and follows all of its gyri and sulci. The cerebral blood vessels enter the brain from its surface by perforating the pia mater. Except for the capillaries, all such vessels are accompanied for a short distance by a pial sheath, and thereafter by a glial membrane that separates them from the neuropil. The perivascular space enclosed by this membrane (*Virchow–Robin space*) contains cerebrospinal fluid. The *choroid plexus* of the cerebral ventricles, which secretes the cerebrospinal fluid, is formed by an infolding of pial blood vessels (*tela choroidea*) covered by a layer of ventricular epithelium (*ependyma*).

Arachnoid Membrane

The dura mater is closely apposed to the arachnoid membrane; the virtual space between them (subdural space) contains capillaries and bridging veins. Between the arachnoid membrane and the pia mater lies the *sub-arachnoid space*, which is filled with cerebrospinal fluid and is spanned by a network of delicate trabecular fibers.



Cerebral Ventricles and Cisterns

The fluid-filled cerebral ventricles constitute the inner CSF space. Each of the two lateral ventricles communicates with the third ventricle through the interventricular foramen of Monro (one on each side). Fluid passes from the third ventricle through the cerebral aqueduct (of Sylvius) into the fourth ventricle, and thence through the single midline foramen (of Magendie) and paired lateral foramina (of Luschka) into the subarachnoid space (outer CSF space). Dilatations of the subarachnoid space are called *cisterns*. The *cerebellomedullary* cistern (cisterna magna) lies between the posterior surface of the medulla and the undersurface of the cerebellum. The cerebellopontine cistern occupies the cerebellopontine angle. The ambient cistern lies lateral to the cerebral peduncle and contains the posterior cerebral and superior cerebellar arteries, the basal vein, and the trochlear nerve. The interpeduncular cistern lies in the midline between the cerebral peduncles and contains the oculomotor nerves, the bifurcation of the basilar artery, and the origins of the superior cerebellar and posterior cerebral arteries: anterior to it is the chiasmatic cistern, which surrounds the optic chiasm and the pituitary stalk. The portion of the subarachnoid space extending from the foramen magnum to the dorsum sellae is collectively termed the posterior cistern.

Cerebrospinal Fluid (CSF)

The CSF, a clear and colorless ultrafiltrate of blood plasma, is mainly produced in the choroid plexus of the cerebral ventricles and in the capillaries of the brain. It normally contains no red blood cells and at most 4 white blood cells/ul. Its functions are both physical (compensation for volume changes, buffering and equal distribution of intracranial pressure despite variation in venous and arterial blood pressure) and metabolic (transport of nutrients and hormones into the brain, and of waste products out of it). The total CSF volume in the adult is ca. 150 ml, of which ca. 30 ml is in the spinal subarachnoid space. Some 500 ml of cerebrospinal fluid is produced per day, corresponding to a flow of ca. 20 ml/h. The normal pulsation of CSF reflects brain pulsation due to changes in cerebral venous and arterial volume, respiration, and head movements. A Valsalva maneuver increases the CSF pressure.

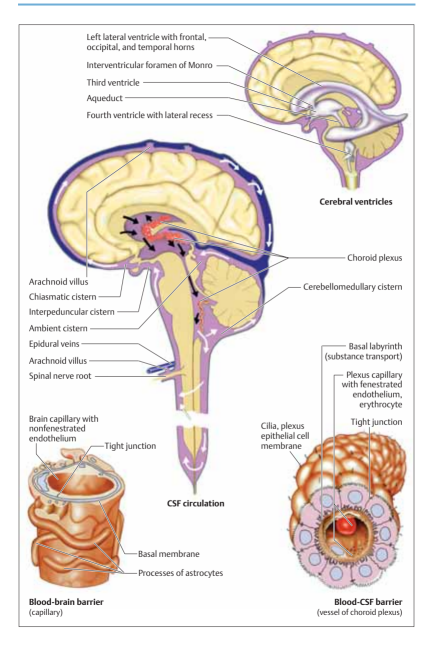
CSF circulation. CSF formed in the choroid plexus flows through the ventricular system and through the foramina of Magendie and Luschka into the basal cisterns. It then circulates further into the spinal subarachnoid space, over the surfaces of the cerebellum and cerebrum, eventually reaching the sites of CSF absorption. It is mainly absorbed through the arachnoid villi (arachnoid granulations, pacchionian corpuscles), which are most abundant along the superior sagittal sinus but are also found at spinal levels. CSF drains through the arachnoid villi in one direction, from the subarachnoid space to the venous compartment, by a valve mechanism. This so-called bulk flow is apparently achieved with the aid of pinocytotic vacuoles that transport the CSF, and all substances dissolved in it, in ladlelike fashion. At the same time, CSF diffuses into the brain tissue adjacent to the CSF space and is absorbed by the capillaries.

The Blood-CSF and Blood-Brain Barriers

These "barriers" are not to be conceived of as impenetrable; under normal conditions, all plasma proteins pass into the CSF. The larger the protein molecule, however, the longer it takes to reach the CSF, and the steeper the plasma/CSF concentration gradient. The term *blood-brain barrier* (BBB) is a collective term for all barriers lying between the plasma and the neuropil, one of which is the *blood-CSF barrier* (BCB). Disease processes often alter the permeability of the BBB, but very rarely that of the BCB.

Morphologically, the BCB is formed by the choroid epithelium, while the BBB is formed by the tight junction (zonula occludens) of capillary endothelial cells. Up to half of all cerebral capillaries have a tubular structure, i.e., they have no connecting interstices. Physiologically, the system of barriers enables the regulation of the osmolarity of brain tissue and CSF and, thereby, the intracranial pressure and volume. Biochemically, the BCB is permeable to water-soluble substances (e.g., plasma proteins) but not to liposoluble substances such as anesthetics, psychoactive drugs, and analgesics. The BBB, on the other hand, is generally permeable to liposoluble substances (of molecular weight less than 500 daltons) but not to water-soluble substances.

Cerebrospinal Fluid



Blood is pumped from the left ventricle of the heart to the aortic arch and thence to the common carotid arteries and anterior circulation of the brain (internal carotid, middle crebral, and anterior cerebral arteries), and to the subclavian arteries and posterior circulation of the brain (vertebral, basilar, and posterior cerebral arteries). The *anterior circulation* supplies the eyes, basal ganglia, part of the hypothalamus, the frontal and parietal lobes, and a large portion of the temporal lobes, while the *posterior circulation* supplies the brain stem, cerebellum, inner ear, occipital lobes, the thalamus, part of the hypothalamus, and a smaller portion of the temporal lobes.

Venous blood from the *superficial and deep cerebral veins* (p. 18 ff) drains via the dural venous sinuses into the internal jugular veins and thence into the the superior vena cava and right atrium. The *extracranial and intracranial portions* of the blood supply of the brain as well as that of the spinal cord will be detailed further in the following paragraphs.

Carotid Arteries: Extracranial Portion

The brachiocephalic trunk arises from the aortic arch behind the manubrium of the sternoum and bifurcates at the level of the sternoclavicular joint to form the right subclavian and common carotid arteries. The left common carotid artery (usually adjacent to the brachiocephalic trunk) and subclavian artery arise directly from the aortic arch. The common carotid artery on either side bifurcates at the level of the thyroid cartilage to form the internal and external carotid arteries; these arteries lie parallel and adjacent to each other after the bifurcation, with the external carotid artery lying medial. A dilatation of the common carotid artery at its bifurcation is called the *carotid sinus*.

The external carotid artery gives off the superior thyroid, lingual, facial, and maxillary arteries anteriorly, the ascending pharyngeal artery medially, and the occipital and posterior auricular arteries posteriorly. The maxillary and superficial temporal arteries are its terminal branches. The *middle meningeal artery* is an important branch of the maxillary artery.

The *internal carotid artery* gives off no extracranial branches. Its cervical portion runs lateral or dorsolateral to the external carotid artery, then dorsomedially along the wall of the pharynx (parapharyngeal space) in front of the transverse processes of the first three cervical vertebrae, and finally curves medially toward the carotid foramen.

Carotid Arteries: Intracranial Portion

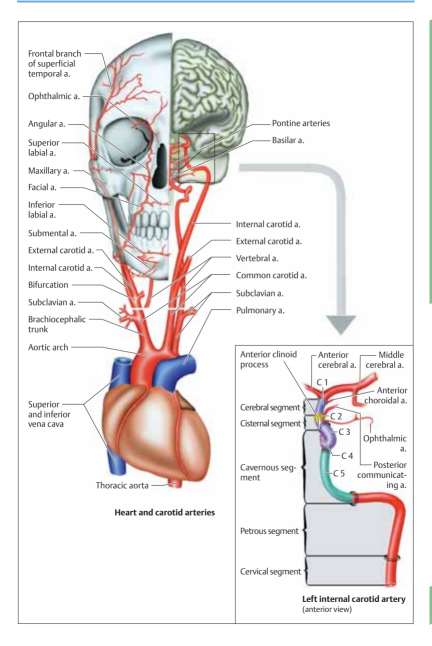
The internal carotid artery (ICA) passes through the base of the skull in the carotid canal, which lies within the petrous part of the temporal bone. It runs upward about 1 cm. then turns anteromedially and courses toward the petrous apex, where it emerges from the temporal bone to enter the cavernous sinus. Within the sinus, the ICA runs along the lateral surface of the body of the sphenoid bone (C5 segment of the ICA), then turns anteriorly and passes lateral to the sella turcica along the lateral wall of the sphenoid bone (segment C4). It then bends sharply back on itself under the root of the anterior clinoid process, so that it points posteriorly (segment C3, carotid bend). After emerging from the cavernous sinus, it penetrates the dura mater medial to the anterior clinoid process and passes under the optic nerve (cisternal segment, segment C2). It then ascends in the subarachnoid space (segment C1) till it reaches the circle of Willis. the site of its terminal bifurcation. Segments C3, C4, and C5 of the ICA constitute its infraclinoid segment, segments C1 and C2 its supraclinoid segment. Segments C2, C3, and C4 together make up the carotid siphon.

The *ophthalmic artery* arises from the carotid bend and runs in the optic canal inferior to the optic nerve. One of its ocular branches, the *central retinal artery*, passes together with the optic nerve to the retina, where it can be seen by ophthalmoscopy.

Medial to the clinoid process, the *posterior communicating artery* arises from the posterior wall of the internal carotid artery, passes posteriorly in proximity to the oculomotor nerve, and then joins the posterior cerebral artery.

The anterior choroidal artery usually arises from the ICA and rarely from the middle cerebral artery. It crosses under the optic tract, passes laterally to the crus cerebri and lateral geniculate body, and enters the inferior horn of the lateral ventricle, where it joins the tela choroidea.

Carotid Arteries



The anterior and middle cerebral arteries are the terminal branches of the internal carotid artery. They originate at the ICA bifurcation, located in the circle of Willis at the level of the anterior clinoid process, between the optic chiasm and the temporal pole.

Anterior Cerebral Artery (ACA)

The ACA is the more medial of the two arteries arising from the ICA bifurcation. It ascends lateral to the anterior clinoid process and past the the optic nerve and optic chiasm, giving off a small branch, the anterior communicating artery (ACommA), which crosses the midline to join the contralateral ACA. The segment of ACA proximal to the origin of the ACommA is its precommunicating segment (segment A1). The A1 segments on either side and the ACommA together form the anterior half of the circle of Willis. Segment A1 gives off an average of eight basal perforating arteries that enter the brain through the anterior perforated substance. The recurrent artery of Heubner arises from the ACA near the origin of the ACommA, either from the distal part of A1 or from the proximal part of A2.

The postcommunicating segment of the ACA (segments A2 to A5) ascends between the frontal lobes and runs toward the occiput in the interhemispheric fissure, along the corpus callosum and below the free border of the falx cerebri, as the pericallosal artery. Segment A2, which usually gives off the frontopolar artery, ends where the artery turns forward to become apposed to the genu of the corpus callosum; segment A3 is the frontally convex arch of the vessel along the genu. The A4 and A5 segments run roughly horizontally over the callosal surface and give off supracallosal branches that run in a posterior direction.

Distribution. The basal perforating arteries arising from A1 supply the ventral hypothalamus and a portion of the pituitary stalk. Heubner's artery supplies the head of the caudate nucleus, the rostral four-fifths of the putamen, the globus pallidus, and the internal capsule. The blood supply of the inferior portion of the genu of the corpus callosum, and of the olfactory bulb, tract, and trigone, is variable.

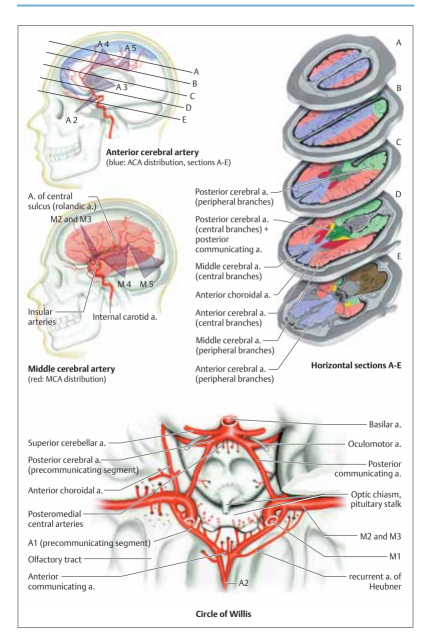
The ACommA gives off a few small branches (anteromedial central branches) to the hypothalamus. Branches from the postcommunicating segment of the ACA supply the inferior surface of the frontal lobe (frontobasilar artery), the medial and parasagittal surfaces of the frontal lobe (callosomarginal artery), the paracentral lobule (paracentral artery), the medial and parasagittal surfaces of the parietal lobe (precuneal artery), and the cortex in the region of the parieto-occipital sulcus (parieto-occipital artery).

Middle Cerebral Artery (MCA)

The MCA is the more lateral of the two arteries arising from the ICA bifurcation. Its first segment (M1, sphenoidal segment) follows the anterior clinoid process for a distance of 1 to 2 cm. The MCA then turns laterally to enter the depths of the Sylvian fissure (i.e., the Sylvian cistern), where it lies on the surface of the insula and gives off branches to it (M2, insular segment). It bends back sharply to travel along the surface of the operculum (M3, opercular segment) and then finally emerges through the Sylvian fissure onto the lateral convexity of the brain (M4 and M5, terminal segments).

Distribution. Small branches of M1 (the thalamostriate and lenticulostriate arteries) supply the basal ganglia, the claustrum, and the internal, external, and extreme capsules. M2 and M3 branches supply the insula (insular arteries), lateral portions of the orbital and inferior frontal gyri (frontobasal artery), and the temporal operculum, including the transverse gyrus of Heschl (temporal arteries). M4 and M5 branches supply most of the cortex of the lateral cerebral convexity, including portions of the frontal lobe (arteries of the precentral and triangular sulci), the parietal lobe (anterior and posterior parietal arteries), and the temporal lobe (arteries of central and postcentral sulci). In particular, important cortical areas supplied by M4 and M5 branches include the primary motor and sensory areas (precentral and postcentral gyri) and the language areas of Broca and Wernicke.

Anterior Circulation of the Brain



Extracranial Portion

The vertebral artery arises from the arch of the subclavian artery at a point designated V0. The prevertebral or V1 segment extends from V0 to the foramen transversarium of the transverse process of C6. The transversarial or V2 segment passes vertically through the foramina transversaria of C6 through C2, accompanied by venous plexuses and sympathetic nerves derived from the cervical ganglia. It gives off branches to the cervical nerves, vertebrae and intervertebral joints, neck muscles, and cervical spinal cord. Often, a prominent branch at the C5 level anastomoses with the anterior spinal artery. The V3 segment, also called the atlas (C1) loop, runs laterally and then vertically to the foramen transversarium of C1, which it passes through, winds medially along the lateral mass of C1, pierces the posterior atlanto-occipital membrane behind the atlanto-occipital joint, and then enters the dura mater and arachnoid membrane at the level of the foramen magnum. The two vertebral arteries are unequal in size in about 75% of persons, and one of them is extremely narrow (hypoplastic) in about 10%, usually on the right side.

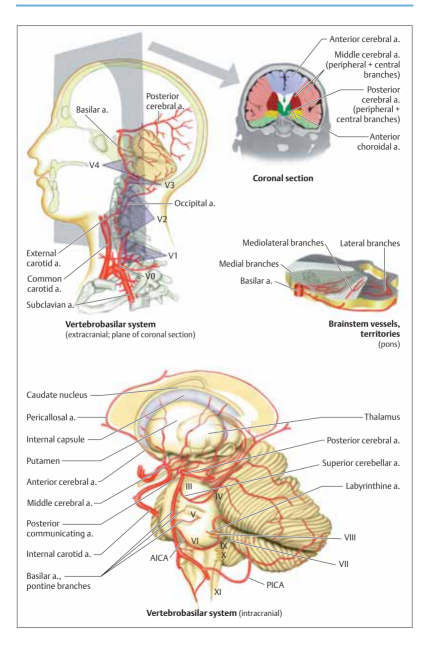
Intracranial Portion

The V4 segment of the vertebral artery lies entirely within the subarachnoid space. It terminates at the junction of the two vertebral arteries to form the basilar artery, at the level of the lower border of the pons. Proximal to the junction, each vertebral artery gives off a mediobasal branch: these two branches run for ca. 2 cm and then unite in the midline to form a single anterior spinal artery, which descends along the anterior surface of the medulla and spinal cord (see p. 23). The posterior inferior cerebellar artery (PICA), which originates from the V4 segment at a highly variable level, curves around the inferior olive and extends dorsally through the root filaments of the accessory nerve. It then ascends behind the fibers of the hypoglossus and vagus nerves, forms a loop on the posterior wall of the fourth ventricle, and gives off terminal branches to the inferior surface of the cerebellar hemisphere, the tonsils, and the vermis. It provides most of the blood supply to the dorsolateral medulla and the posteroinferior surface of the cerebellum. The *posterior spinal artery* (there is one on each side) arises from either the vertebral artery or the PICA.

The *basilar artery* runs in the prepontine cistern along the entire length of the pons and then bifurcates to form the posterior cerebral arteries. Its inferior portion is closely related to the abducens nerves, its superior portion to the oculomotor nerves. Its paramedian, short circumferential, and long circumferential branches supply the pons and the superior and middle cerebellar peduncles.

The anterior inferior cerebellar artery (AICA) arises from the lower third of the basilar artery. It runs laterally and caudally toward the cerebellopontine angle, passes near the internal acoustic meatus, and reaches the flocculus, where it gives off terminal branches that supply the anteroinferior portion of the cerebellar cortex and part of the cerebellar nuclei. The AICA lies basal to the abducens nerve and ventromedial to the facial and auditory nerves in the cerebellopontine cistern. It often gives rise to a labyrinthine branch that enters the internal acoustic meatus.

The superior cerebellar arteries (SCA) of both sides originate from the basilar trunk just below its bifurcation. Each SCA travels through the perimesencephalic cistern dorsal to the oculomotor nerve, curves around the cerebral peduncle caudal and medial to the trochlear nerve, and then enters the ambient cistern, where it gives off its terminal branches. The SCA supplies the upper pons, part of the mid brain, the upper surface of the cerebellar hemispheres, the upper portion of the vermis, and the cerebellar nuclei.



Posterior Cerebral Artery (PCA)

The precommunicating segment of the PCA (P1) extends from the basilar bifurcation to the origin of the posterior communicating artery (PCommA). Its course lies within the interpeduncular cistern, which is demarcated by the clivus and the two cerebral peduncles. The oculomotor nerve, after its emergence from the brain stem, runs between the PCA and the superior cerebellar artery. The postcommunicating segment (P2) curves laterally and backward around the crus cerebri and reaches the posterior surface of the midbrain at an intercollicular level.

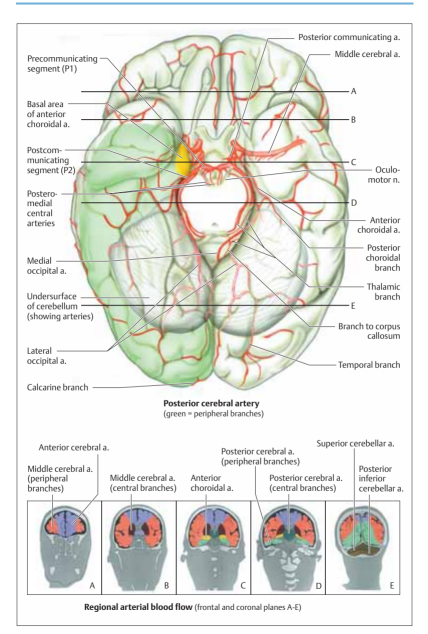
The precommunicating and postcommunicating segments are together referred to as the *pars circularis* of the PCA. (Alternatively, the pars circularis may be divided into three segments interpeduncular, ambient, and quadrigeminal named after the cisterns they traverse.)

Distal to the pars circularis of the PCA is the pars terminalis, which divides above the tentorium and caudal to the lateral geniculate body to form its terminal branches, the medial and lateral occipital arteries.

Pars circularis. The precommunicating segment gives off fine branches (posteromedial central arteries) that pierce the interpeduncular perforated substance to supply the anterior thalamus, the wall of the third ventricle, and the globus pallidus. The postcommunicating segment gives off fine branches (posterolateral central arteries) to the cerebral peduncles, the posterior portion of the thalamus, the colliculi of the mid brain, the medial geniculate body, and the pineal body. Further branches supply the posterior portion of the thalamus (thalamic branches), the cerebral peduncle (peduncular branches), and the lateral geniculate body and choroid plexus of the third and lateral ventricles (posterior choroidal branches).

Pars terminalis. Of the two terminal branches of this terminal portion of the PCA, the lateral occipital artery (together with its temporal branches) supplies the uncus, the hippocampal gyrus, and the undersurface of the occipital lobe. The medial occipital artery passes under the splenium of the corpus callosum, giving off branches that supply it (dorsal branch to the corpus callosum) as well as the cuneus and pre-

cuneus (parieto-occipital branch), the striate cortex (calcarine branch), and the medial surfaces of the occipital and temporal lobes (occipitotemporal and temporal banches), including the parasagittal portion of the occipital lobe.



Cerebral Veins

The superficial cerebral veins (cortical veins) carry blood from the outer 1–2 cm of the brain surface to large drainage channels such as the superior and inferior sagittal sinuses, the great cerebral vein of Galen, the straight sinus, and the tentorial veins. Thus, the cerebellar veins drain blood from the cerebellar surface into the great cerebral vein, straight sinus, and transverse sinuses. The *deep cerebral veins* (central veins) drain blood from the inner regions of the brain (hemispheric white matter, basal ganglia, corpus callosum, choroid plexus) and from a few cortical areas as well.

Superficial cerebral veins (cortical veins). The superficial cerebral veins are classified by their location as prefrontal, frontal, parietal, and occipital. Except for the occipital veins, which empty into the transverse sinus, these veins all travel over the cerebral convexity to join the superior sagittal sinus. They are termed bridging veins at their distal end, where they pierce the arachnoid membrane and bridge the subarachnoid space to join the sinus. The superficial middle cerebral vein (not shown) usually follows the posterior ramus of the Sylvian fissure and the fissure itself to the cavernous sinus. The inferior cerebral veins drain into the cavernous sinus, superior petrosal sinus, and transverse sinus. The superior cerebral veins drain into the superior sagittal sinus.

Deep cerebral veins (central veins). The internal cerebral vein arises bilaterally at the level of the interventricular foramen (of Monro). It traverses the transverse cerebral fissure to a point just inferior to the splenium of the corpus callosum. The venous angle at its junction with the superior thalamostriate vein can be seen in a laterally projected angiogram. The two internal cerebral veins join under the splenium to form the great cerebral vein (of Galen), which receives the basal vein (of Rosenthal) and then empties into the straight sinus at the anterior tentorial edge at the level of the quadrigeminal plate. The basal vein of Rosenthal is formed by the union of the anterior cerebral vein, the deep middle cerebral vein, and the striate veins. It passes posteromedial to the optic tract, curves around the cerebral peduncle, and empties into the internal vein or

the great cerebral vein posterior to the brain stem.

Posterior fossa. The anterior, middle, and posterior veins of the posterior fossa drain into the great cerebral vein, the petrosal vein, and the tentorial and straight sinuses, respectively.

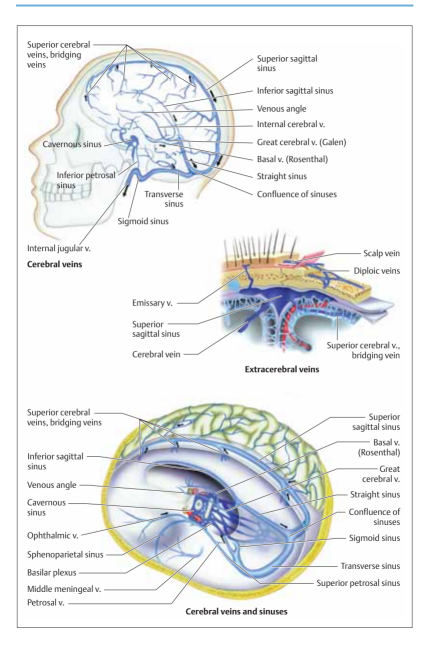
Extracerebral Veins

The extracerebral veins—most prominently, the dural venous sinuses—drain venous blood from the brain into the sigmoid sinuses and jugular veins.

The *diploic veins* drain into the extracranial veins of the scalp and the intracranial veins (dural venous sinuses).

The *emissary veins* connect the sinuses, diploic veins, and superficial veins of the skull. Infections sometimes travel along the emissary veins from the extracranial to the intracranial compartment.

The veins of the brain empty into the superior and inferior groups of dural venous sinuses. The sinuses of the superior group (the superior and inferior sagittal, straight, and occipital sinuses) join at the confluence of the sinuses (torcular Herophili), which drains into both transverse sinuses and thence into the sigmoid sinuses and internal jugular veins. The sinuses of the inferior group (superior and inferior petrosal sinuses) join at the cavernous sinus, which drains into the sigmoid sinus and internal jugular vein via the inferior petrosal sinus, or into the internal vertebral plexus via the basilar plexus.



Craniocervical Veins

Anastomotic channels connect the cutaneous veins of the two sides of the head. Venous blood from the facial, temporal, and frontal regions drains into the facial and retromandibular veins and thence into the internal jugular vein. Some blood from the forehead drains via the nasofrontal, angular, and superior ophthalmic veins into the cavernous sinus. The occipital vein carries blood from the posterior portion of the scalp into the deep cervical vein and thence into the external jugular vein. Blood from the jugular veins continues to the brachiocephalic vein, superior vena cava, and right atrium. The venous channels in the spinal canal and the transcranial emissary veins play no more than a minor role in venous drainage. The pterygoid plexus links the cavernous sinus, the facial vein, and the internal jugular vein.

The numerous anastomoses between the extracranial and intracranial venous systems provide a pathway for the spread of infection from the scalp or face to the intracranial compartment. For example, periorbital infection may extend inward and produce septic thrombosis of the cavernous sinus.

Cranial Veins

The facial vein drains the venous blood from the face and anterior portion of the scalp. It begins at the inner canthus as the angular vein and communicates with the cavernous sinus via the superior ophthalmic vein. Below the angle of the mandible, it merges with the retromandibular vein and branches of the superior thyroid and superior laryngeal veins. It then drains into the internal jugular vein in the carotid triangle. The veins of the temporal region, external ear, temporomandibular joint, and lateral aspect of the face join in front of the ear to form the retromandibular vein, which either joins the facial vein or drains directly into the internal jugular vein. Its upper portion gives off a prominent dorsocaudal branch that joins the posterior auricular vein over the sternocleidomastoid muscle to communicate with the external jugular vein. Venous blood from the posterior portion of the scalp and the mastoid and occipital emissary veins drains into the occipital vein, which anastomoses with the occipital venous plexus and finally drains into the external jugular vein.

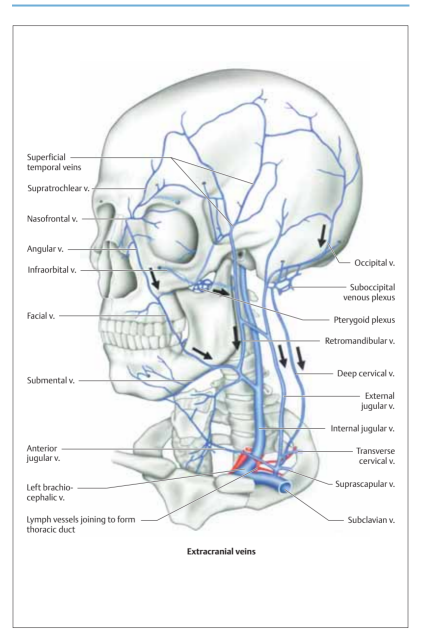
The *pterygoid plexus* lies between the temporalis, medial pterygoid, and lateral pterygoid muscles and receives blood from deep portions of the face, the external ear, the parotid gland, and the cavernous sinus, which it carries by way of the maxillary and retromandibular veins to the internal jugular vein.

Cervical Veins

The *deep cervical vein* originates from the occipital vein and suboccipital plexus. It follows the course of the deep cervical artery and vertebral artery to arrive at the brachiocephalic vein, which it joins.

The vertebral vein, which also originates from the occipital vein and suboccipital plexus, envelops the vertebral artery like a net and accompanies it through the foramina transversaria of the cervical vertebrae, collecting blood along the way from the cervical spinal cord, meninges, and deep neck muscles through the vertebral venous plexus, and finally joining the brachiocephalic vein.

Extracranial Veins



Arteries

Most of the blood supply of the spinal cord is supplied by the segmental spinal arteries, while relatively little comes from the vertebral arteries via the anterior and posterior spinal arteries. The segmental and spinal arteries are linked by numerous anastomoses.

Segmental arteries. The vertebral, ascending cervical, and deep cervical arteries give off cervical segmental branches; the thoracic and abdominal aorta give off thoracolumbar segmental branches via the posterior intercostal and lumbar arteries.

The segmental arteries give off radicular branches that enter the intervertebral foramen and supply the anterior and posterior roots and spinal ganglion of the corresponding level. The spinal cord itself is supplied by unpaired medullary arteries that originate from segmental arteries. The anatomy of these medullary arteries is variable; they usually have 5 to 8 larger ventral and dorsal branches that join up with the anterior and posterior spinal arteries. Often there is a single large radicular branch on one side, the great radicular artery (of Adamkiewicz), that supplies the entire lower twothirds of the spinal cord. It usually enters the spinal canal in the lower thoracic region on the left side.

Spinal arteries. The spinal arteries run longitudinally down the spinal cord and arise from the vertebral artery (p. 14). The unpaired anterior spinal artery lies in the anterior median fissure of the spinal cord and supplies blood to the anterior two-thirds of the cord. The artery's diameter steadily increases below the T2 level. The two posterior spinal arteries supply the dorsal columns and all but the base of the dorsal horns bilaterally. Numerous anastomoses of the spinal arteries produce a vasocorona around the spinal cord. The depth of the spinal cord is supplied by these arteries penetrating it from its outer surface and by branches of the anterior spinal artery penetrating it from the anterior median fissure (sulcocommissural arteries).

Spinal Veins

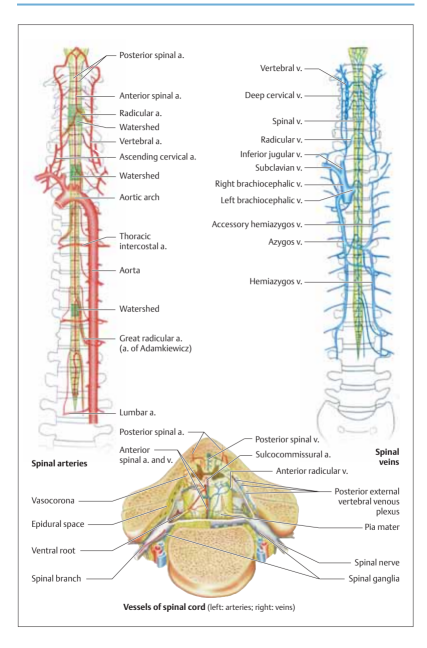
Blood from within the spinal cord travels through the intramedullary veins, whose anat-

omy is variable, to the anterior and posterior spinal veins, which form a reticulated network in the pia mater around the circumference of the cord and down its length. The anterior spinal vein drains the anterior two-thirds of the gray matter, while the posterior and lateral spinal veins drain the rest of the spinal cord. These vessels empty by way of the radicular veins into the external and internal vertebral venous plexuses, groups of valveless veins that extend from the coccvx to the base of the skull and communicate with the dural venous sinuses via the suboccipital veins. Venous blood from the cervical spine drains by way of the vertebral and deep cervical veins into the superior vena cava; from the thoracic and lumbar spine, by way of the posterior intercostal and lumbar veins into the azygos and hemiazygos veins; from the sacrum, by way of the median and lateral sacral veins into the common iliac vein

Watershed Zones

Because blood can flow either upward or downward in the anterior and posterior spinal arteries, the tissue at greatest risk of hypoperfusion is that located at a border zone between the distributions of two adjacent supplying arteries ("watershed zone"). Such vulnerable zones are found in the cervical, upper thoracic, and lower thoracic regions (ca. C4, T3–T4, and T8–T9).

Spinal Circulation



Cortical Structures

Different areas of the *cerebral cortex* (neocortex) may be distinguished from one another by their histological features and neuroanatomical connections. *Brodmann's* numbering scheme for cortical areas has been used for many years and will be introduced in this section.

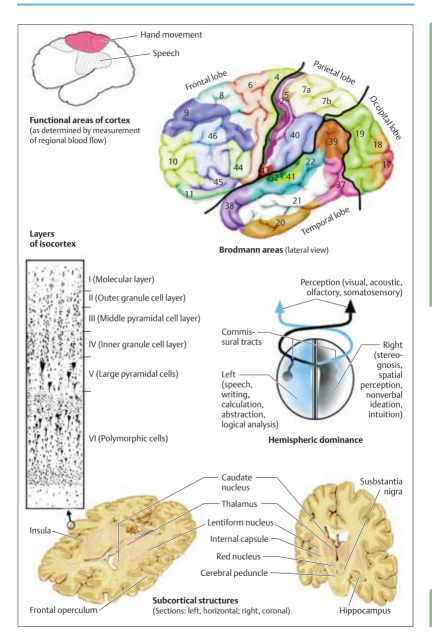
Projection areas. By following the course of axons entering and leaving a given cortical area, one may determine the other structures to which it is connected by afferent and efferent pathways. The primary projection areas are those that receive most of their sensory impulses directly from the thalamic relay nuclei (primary somatosensory cortex; Brodman areas 1, 2, 3), the visual (area 17), or the auditory (areas 41, 42) pathways. The primary motor cortex (area 4) sends motor impulses directly down the pyramidal pathway to somatic motor neurons within brainstem and the spinal cord. The primary projection areas are somatotopically organized and serve the contralateral half of the body. Proceeding outward along the cortical surface from the primary projection areas, one encounters the secondary projection areas (motor, areas 6, 8, 44; sensory, areas 5, 7a, 40; visual, area 18; auditory, area 42), which subserve higher functions of coordination and information processing, and the tertiary projection areas (motor, areas 9, 10, 11; sensory, areas 7b, 39; visual, areas 19, 20, 21; auditory, area 22), which are responsible for complex functions such as voluntary movement, spatial organization of sensory input, cognition, memory, language, and emotion. The two hemispheres are connected by commissural fibers, which enable bihemispheric coordination of function. The most important commissural tract is the corpus callosum; because many tasks are performed primarily by one of the two hemispheres (cerebral dominance), interruption of the corpus callosum can produce various disconnection syndromes. Total callosal transection causes splitbrain syndrome, in which the patient cannot name an object felt by the left hand when the eyes are closed, or one seen in the left visual hemifield (tactile and optic anomia), and cannot read words projected into the left visual hemifield (left hemialexia), write with the left hand (left hemiagraphia), or make pantomimic movements with the left hand (left hemiapraxia). Anterior callosal lesions cause *alien hand syndrome* (diagonistic apraxia), in which the patient cannot coordinate the movements of the two hands. Disconnection syndromes are usually not seen in persons with congenital absence (agenesis) of the corpus callosum.

Cytoarchitecture. Most of the cerebral cortex consists of *isocortex*, which has six distinct cytoarchitectural layers. The Brodmann classification of cortical areas is based on distinguishing histological features of adjacent areas of isocortex.

Functional areas. The functional organization of the cerebral cortex can be studied with various techniques: direct electrical stimulation of the cortex during neurosurgical procedures. measurement of cortical electrical cortical activity (electroencephalography and evoked potentials), and measurement of regional cerebral blood flow and metabolic activity. Highly specialized areas for particular functions are found in many different parts of the brain. A lesion in one such area may produce a severe functional deficit, though partial or total recovery often occurs because adjacent uninjured areas may take over some of the function of the lost brain tissue. (The extent to which actual brain regeneration may aid functional recovery is currently unclear.) The specific anatomic patterns of functional localization in the brain are the key to understanding much of clinical neurology.

Subcortical Structures

The subcortical structures include the basal ganglia, thalamus, subthalamic nucleus, hypothalamus, red nucleus, substantia nigra, cerebellum, and brain stem, and their nerve pathways. These structures perform many different kinds of complex information processing and are anatomically and functionally interconnected with the cerebral cortex. Subcortical lesions may produce symptoms and signs resembling those of cortical lesions; special diagnostic studies may be needed for their precise localization.



25

Brain Stem

The brain stem consists of the midbrain (mesencephalon), pons, and medulla. It contains the nuclei of the cranial nerves and ascending and descending tracts running to and from the brain, cerebellum, and spinal cord. It also contains autonomic centers that regulate cardiovascular function, breathing, and eating behavior as well as acoustic and vestibular relay nuclei. The flow of information along afferent and efferent pathways is regulated by reflex systems.

Nerve Pathways

All motor (p. 44) and sensory projection systems (p. 104) pass through the brain stem and communicate with its intrinsic structures at various sites. The central sympathetic pathway (p. 90) originates in the hypothalamus.

Reticular Formation

The reticular formation (RF) is a network of nuclei and interconnecting fibers that is anatomically intertwined with the cranial nerve nuclei and other fiber tracts of the brain stem. Different parts of the reticular formation perform different functions. The reticular activating system (RAS) provides the anatomical and physiological basis for wakeful consciousness (p. 116). The medullary RF contains the vital centers controlling the heartbeat, breathing, and circulation as well as reflex centers for swallowing and vomiting. The pontine RF contains centers for coordination of acoustic, vestibular, respiratory, and cardiovascular processes. The midbrain RF contains centers subserving visuospatial orientation and eating behavior (chewing, sucking, licking).

Reflex Systems (pp. 118ff)

Pupillary light reflex. The Edinger–Westphal nucleus in the midbrain, which is adjacent to the oculomotor nucleus, provides the efferent arm of the reflex loop (p. 90; examination, p. 92.) Vestibulo-ocular reflex (VOR, p. 84). The vestibular nuclei receive their main input from the labyrinthine semicircular canals and collateral input from the cerebellar nuclei; their output is conveyed to the extraocular muscles through the medial longitudinal fasciculus, and to the

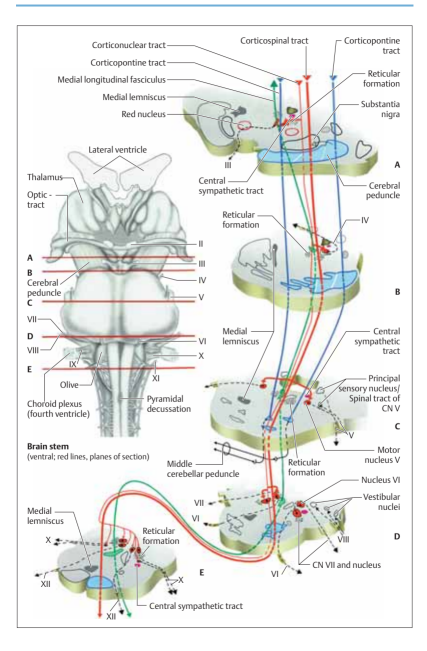
spinal cord through the vestibulospinal tract. Examination: Suppression of visual fixation: the subject extends his arms and stares at his thumbs while spinning on a swivel chair. Nystagmus does not occur in normal subjects. Oculocephalic reflex (doll's eyes phenomenon): Horizontal or vertical passive rotation of the subject's head causes the eyes to rotate in the opposite direction; normally suppressible by awake persons, this reflex is seen in patients with impaired consciousness but preserved vestibular function. Caloric testing: The examiner first confirms that the patient's eardrums are intact, then instills cold water in the external auditory canal with the head elevated at a 30° angle (which inactivates the ipsilateral horizontal semicircular canal). This normally causes nystagmus in the contralateral direction, i.e., slow ipsilateral conjugate deviation of the eyes, followed by a quick jerk to the other side.

Corneal reflex. Afferent arm, CN V/1; efferent arm, CN VII, which innervates the orbicularis oculi muscle. *Examination*: Touching the cornea from the side while the subject looks forward evokes blinking. The reflex can also be assessed by electromyography (EMG).

Pharyngeal (gag) reflex. Afferent arm, mainly CN IX, X, and V/2; efferent arm, CN IX and X. The gag reflex may be absent in normal persons. Examination: Touching the soft palate or back of the pharynx evokes pharyngeal muscle contraction. Cough reflex. Afferent arm, CN IX and X; efferent arm, via the solitary tract to the diaphragm and other participating muscle groups. Examination: Tested in intubated patients with endotracheal suction (tracheal reflex).

Masseter (jaw jerk) reflex. *Afferent arm*, probably CN V/3; *efferent arm*, CN V. *Examination*: Tapping the chin evokes jaw closure.

Acoustic reflex (p. 68). Afferent arm, projections of the cochlear nuclei to the RAS. Examination: Sudden, intense acoustic stimuli evoke a fright reaction including lid closure, startle, turning of the head, and increased alertness.



27

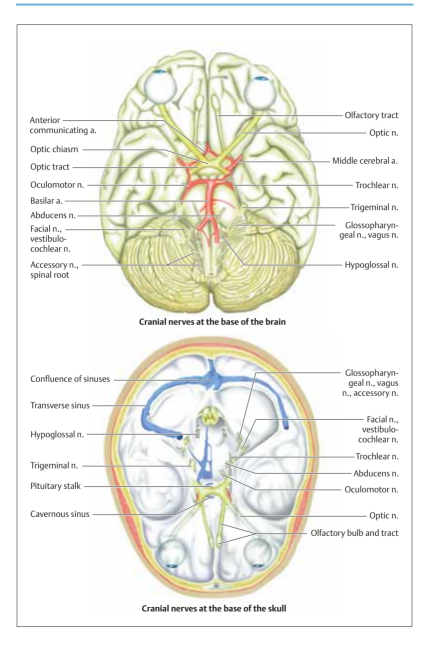
Cranial Nerve Pathways

Cranial Nerve	Origin/Course (see also pp. 70 ff. and 74 ff)		
I	Olfactory nerves \Rightarrow cribriform plate \Rightarrow olfactory bulb \Rightarrow olfactory tract \Rightarrow anterior perforated sub- stance \Rightarrow lateral olfactory stria (\Rightarrow parahippocampal gyrus) and medial olfactory stria (\Rightarrow limbic system)		
II	Retinal ganglion cells \Rightarrow optic disk \Rightarrow optic nerve \Rightarrow orbit \Rightarrow optic canal \Rightarrow optic chiasm \Rightarrow optic tract \Rightarrow lateral geniculate body (\Rightarrow optic radiation \Rightarrow occipital lobe) and superior colliculi (\Rightarrow pretectal area)		
III	Midbrain \Box interpeduncular fossa \Box between superior cerebellar artery and posterior cerebral artery \Box tentorial edge \Box cavernous sinus \Box medial orbital fissure \Box oculomotor nerve, superior division (levator palpebrae and superior rectus muscles) and inferior division (medial and inferior rectus and inferior oblique muscles) or parasympathetic fibers \Box ciliary ganglion		
IV	Midbrain \Rightarrow dorsal brainstem below the inferior colliculi \Rightarrow around the cerebral peduncle \Rightarrow lateral wall of the cavernous sinus \Rightarrow orbital fissure \Rightarrow superior oblique muscle		
V	Pons ⇔ ca. 50 root filaments (sensory root = portio major; motor root = portio minor) ⇒ petrous apex ⇔ through the dura mater ⇔ trigeminal ganglion (V/1 ⇔ orbital fissure; V/2 ⇔ foramen rotundum, V/3 + portio minor ⇔ foramen ovale)		
VI	Posterior margin of pons \Rightarrow up the clivus \Rightarrow through the dura mater \Rightarrow petrous apex \Rightarrow lateral to internal carotid artery in the cavernous sinus \Rightarrow orbital fissure \Rightarrow lateral rectus muscle		
VII	Pons (cerebellopontine angle) above the olive \Rightarrow internal acoustic meatus \Rightarrow petrous pyramid (canal of facial nerve) \Rightarrow geniculum of facial nerve (\Rightarrow nervus intermedius/greater petrosal nerve \Rightarrow gustatory fibers) \Rightarrow medial wall of the tympanic cavity \Rightarrow stylomastoid foramen \Rightarrow muscles of facial expression		
VIII	Lateral to CN VII \Rightarrow vestibular nerve, cochlear nerve		
IX	Medulla \leftrightarrows jugular foramen \rightleftharpoons between carotid artery and internal jugular vein \Leftrightarrow root of the tongue		
Х	Posterolateral sulcus of medulla \Rightarrow jugular foramen \Rightarrow internal organs		
XI	Cranial and spinal roots \Rightarrow trunk of accessory nerve \Rightarrow jugular foramen \Rightarrow muscles		
XII	Medulla ⇔ hypoglossal canal ⇔ tongue muscles		

CN = cranial nerve.

See Table 1, p. 356, for the functions of the cranial nerves.

Cranial Nerves



Spine

The spine (vertebral column) bears the weight of the head, neck, trunk and upper extremities. Its flexibility is greatest in the cervical region, intermediate in the lumbar region, and lowest in the thoracic region. Its uppermost vertebrae (atlas and axis) articulate with the head, and its lowermost portion, the sacrum (which consists of 5 vertebrae fused together), articulates with the pelvis. There are 7 *cervical*, 12 *thoracic* (in British usage *dorsal*), and 5 *lumbar vertebrae*, making a total of 24 above the sacrum. Below the sacrum, the coccyx is composed of 3 to 6 *coccygeal vertebrae*.

Intervertebral Disks

Each pair of adjacent vertebrae is separated by an intervertebral disk. From the third decade of life onward, each disk progressively diminishes in water content, and therefore also in height. Its tensile, fibrous outer ring (annulus fibrosus) connects it with the vertebrae above and below and is held taut by the pressure in the central nucleus pulposus, which varies as a function of the momentary position of the body. The pressure that obtains in the sitting position is double the pressure when the patient stands, but that found in the recumbent position is only onethird as great. The interior of the disk has no nociceptive innervation, in contrast to the periosteum of the vertebral bodies, which is innervated by the meningeal branch of the segmental spinal nerve, as are the intervertebral joint capsules, the posterior longitudinal ligament, the dorsal portion of the annulus fibrosus, the dura mater, and the blood vessels.

Spinal Canal

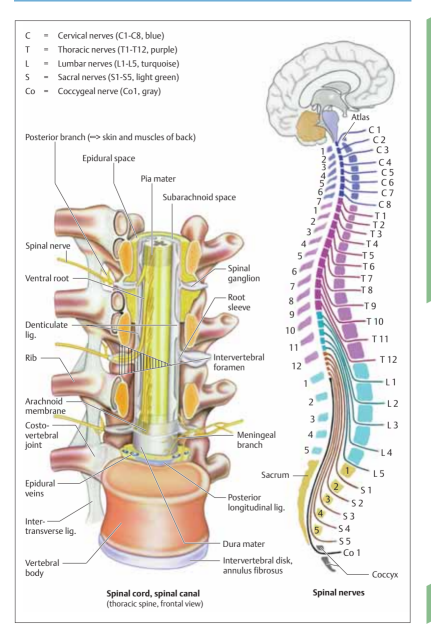
The spinal canal is a tube formed by the vertebral foramina of the vertebral bodies stacked one on top of another; it is bounded anteriorly by the vertebral bodies and posteriorly by the vertebral arches (laminae). Its walls are reinforced by the intervertebral disks and the anterior and posterior longitudinal ligaments. It contains the spinal cord and its meninges, the surrounding fatty and connective tissues, blood vessels, and spinal nerve roots. Its normal sagittal diameter ranges from 12 to 22 mm in the cervical region and from 22 to 25 mm in the lumbar region.

Spinal Cord

Like the brain, the spinal cord is intimately enveloped by the pia mater, which contains numerous nerves and blood vessels; the pia mater merges with the endoneurium of the spinal nerve rootlets and also continues below the spinal cord as the *filum terminale internum*. The weblike spinal arachnoid membrane contains only a few capillaries and no nerves. The denticulate ligament runs between the pia mater and the dura mater and anchors the spinal cord to the dura mater. In lumbar puncture, cerebrospinal fluid is withdrawn from the space between the arachnoid membrane and pia mater (spinal subarachnoid space), which communicates with the subarachnoid space of the brain. The spinal dura mater originates at the edge of the foramen magnum and descends from it to form a tubular covering around the spinal cord. Its lumen ends at the S1-S2 level, where it continues as the filum terminale externum, which attaches to the sacrum, thus anchoring the dura mater inferiorly. The dura mater forms sleeves around the anterior and posterior spinal nerve roots which continue distally, together with the arachnoid membrane, to form the epineurium and perineurium of the spinal nerves. Unlike the cranial dura mater, the spinal dura mater is not directly apposed to the periosteum of the surrounding bone (i.e., the vertebral canal) but is separated from it by the epidural space, which contains fat, loose connective tissue, and valveless venous plexuses (p. 22).

The root filaments (rootlets) that come together to form the ventral and dorsal spinal nerve roots are arranged in longitudinal rows on the lateral surface of the spinal cord on both sides. The ventral root carries only motor fibers, while the dorsal root carries only sensory fibers. (This socalled "law of Bell and Magendie" is actually not wholly true; the ventral root is now known to carry a small number of sensory fibers as well.) The cell bodies of the pseudounipolar sensory neurons are contained in the dorsal root ganglion, a swelling on the dorsal root just proximal to its junction with the ventral root to form the segmental spinal nerve.

Spine and Spinal Cord



The precise region of impaired sensation to light touch and noxious stimuli is an important clue for the clinical localization of spinal cord and peripheral nerve lesions. Reflex abnormalities and autonomic dysfunction are further ones, as discussed below (p. 40, p. 110).

Dermatomes (pp. 34, 36)

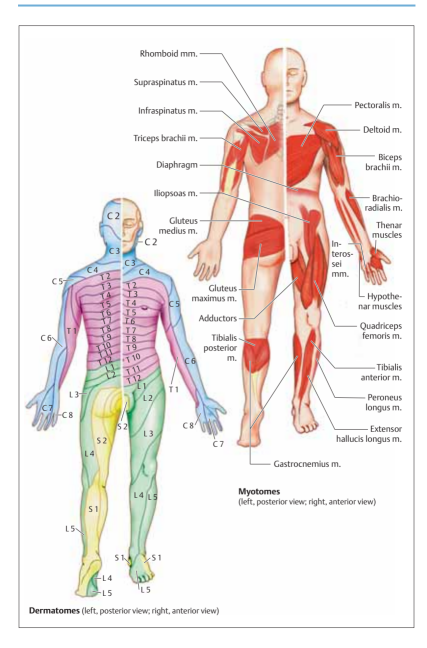
A dermatome is defined as the cutaneous area whose sensory innervation is derived from a single spinal nerve (i.e., dorsal root). The division of the skin into dermatomes reflects the segmental organization of the spinal cord and its associated nerves. Pain dermatomes are narrower, and overlap with each other less, than touch dermatomes (p. 104); thus, the level of a spinal cord lesion causing sensory impairment is easier to determine by pinprick testing than by light touch. (The opposite is true of peripheral nerve lesions.) Radicular pain is pain in the distribution of a spinal nerve root, i.e., in a dermatome; pseudoradicular pain may occupy a bandlike area but cannot be assigned to any particular dermatome. Pseudoradicular pain can be caused by tendomyosis (pain in the muscles that move a particular joint), generalized tendomyopathy or fibromyalgia, facet syndrome (inflammation of the intervertebral joints), myelogelosis (persistent muscle spasm resulting from overexertion), and other conditions. For mnemonic purposes, it is useful to know that the C2 dermatome begins in front of the ear and ends at the occipital hairline; the T1 dermatome comes to the midline of the forearm; the T4 dermatome is at the level of the nipples (which, however, belong to T5); the T10 dermatome includes the navel; the L1 dermatome is in the groin; and the S1 dermatome is at the outer edge of the foot and heel.

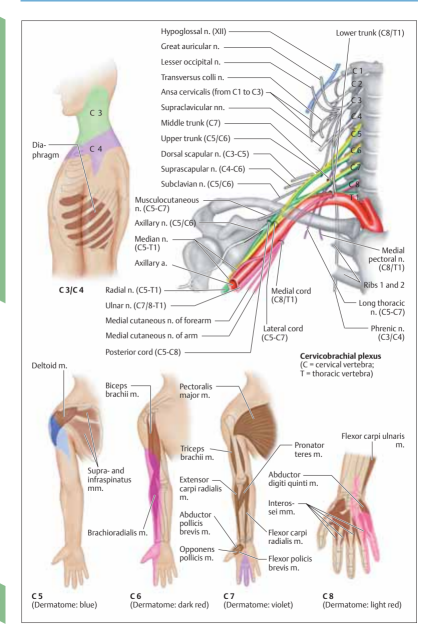
Myotomes

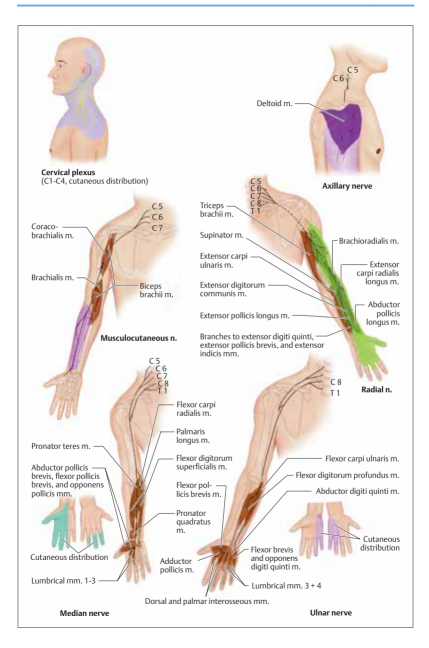
A myotome is defined as the muscular distribution of a single spinal nerve (i.e., ventral root), and is thus the muscular analogue of a cutaneous dermatome. Many muscles are innervated by multiple spinal nerves; only in the paravertebral musculature of the back (erector spinae muscle) is the myotomal pattern clearly segmental (p.31); the nerve supply here is through the dorsal branches of the spinal nerves. Knowledge of the myotomes of each spinal nerve, and of the *segment-indicating muscles* (Table 2, p. 357) in particular, enables the clinical and electromyographic localization of radicular lesions causing motor dysfunction. The segment-indicating muscles are usually innervated by a single spinal nerve, or by two, though there is anatomic variation.

Plexuses (pp. 34, 36) and Peripheral Nerves (pp. 35, 37)

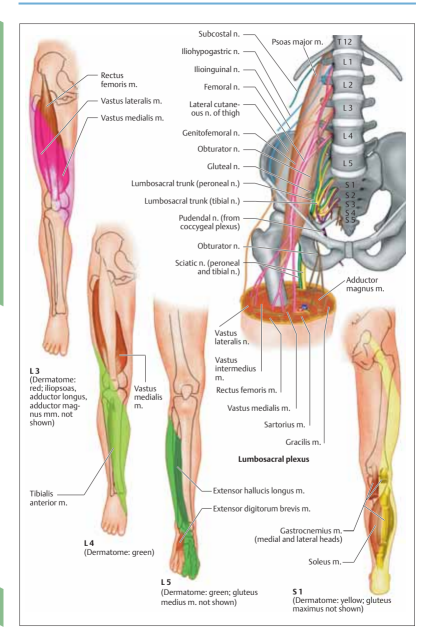
The ventral branches of spinal nerves supplying the limbs join together to form the cervical (C1-C4), brachial (C5-T1), lumbar (T12-L4), and sacral plexuses (L4-S4). The brachial plexus begins as three trunks, the upper (derived from the C5 and C6 roots), middle (C7), and lower (C8, T1). These trunks split into divisions, which recombine to form the lateral (C5–C7), posterior (C5-C8), and medial (C8 and T1) cords (named by their relation to the axillary artery). The cords of the brachial plexus branch into the nerves of the upper limb (p. 35). The nerves of the anterior portion of the lower limb are derived from the lumbar plexus, which lies behind and within the psoas major muscle (p. 37); those of the posterior portion of the lower limb from the sacral plexus. The coccygeal nerve (the last spinal nerve to emerge from the sacral hiatus) joins with the S3-S5 nerves to form the coccygeal plexus, which innervates the coccygeus and the skin over the coccyx and anus (mediates the pain of coccygodynia).



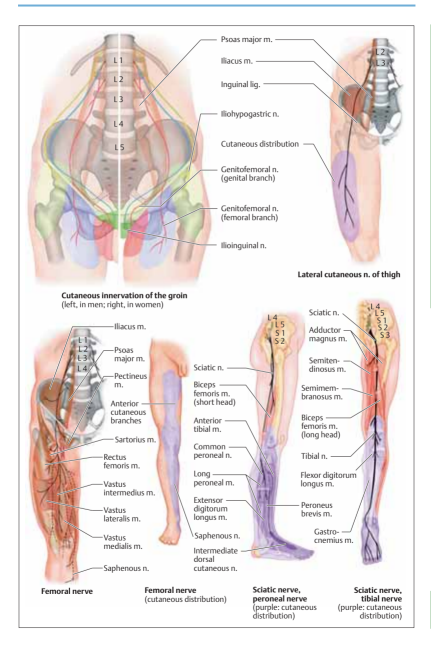




Lumbar Plexus



36



Normal and Abnormal Function of the Nervous System

- Neural Pathways
- Pathophysiology
- Major Syndromes

Reflexes

Reflexes are involuntary and relatively stereotyped responses to specific stimuli. Afferent nerve fibers conduct the impulses generated by activated receptors to neurons in the central nervous system, which fire impulses that are then transmitted through efferent nerve fibers to the cells, muscles, or organs that carry out the reflex response. The pathway as a whole is known as the reflex arc. Receptors are found at the origin of all sensory pathways-in the skin, mucous membranes, muscles, tendons, and periosteum, as well as in the retina, inner ear, olfactory mucosa, and taste buds. A reflex response may involve the somatic musculature or the internal organs. Most reflexes are relatively independent of the state of consciousness. An interruption of the reflex arc at any point weakens or abolishes the reflex. Intrinsic reflexes are those whose receptors and effectors are located in the same organ (e.g., the quadriceps reflex), while the receptors and effectors of extrinsic reflexes are in different organs (e.g., the oculovestibular reflex). Reflexes are important for normal function (e.g., for postural control and goal-directed movement), and an impaired reflex is an important objective finding in clinical neurological examination.

Intrinsic Muscle Reflexes (Phasic Stretch Reflexes, Tendon Reflexes)

Intrinsic muscle reflexes are triggered by stretch receptors within the muscle (annulospiral nerve endings of muscle spindles). The impulses generated at the receptors are conveyed via afferent fast-twitch la fibers to spinal alpha-motor neurons, whose efferent α_1 processes excite the agonistic muscle of an opposing muscle pair. The antagonistic muscle is simultaneously inhibited by spinal interneurons. The resulting muscle contraction relaxes the muscle spindles, thereby stopping impulse generation at the stretch receptors. The spinal reflex arc is also under the influence of higher motor centers.

Abnormal reflex responses imply an abnormality of the musculature, the reflex arc, or higher motor centers. The most important reflexes in clinical diagnosis are the biceps (C5–C6), brachioradialis (C5–C6), triceps (C2–C8), adductor (L2–L4), quadriceps (L2/3–L4), posterior tibial (L5), and Achilles (S1–S2) reflexes.

Extrinsic Reflexes

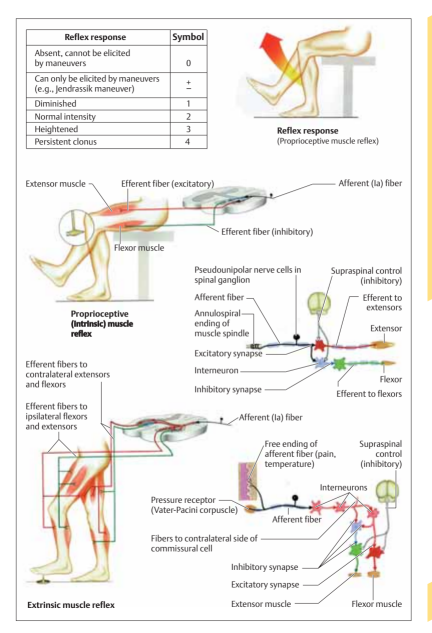
Intrinsic muscle reflexes, discussed above, are monosynaptic, but extrinsic reflexes are polysynaptic: between their afferent and efferent arms lies a chain of spinal interneurons. They may be activated by stimuli of various types, e.g., muscle stretch, touch on the skin (abdominal reflex) or cornea (corneal reflex), mucosal irritation (sneezing), light (eye closure in response to a bright flash), or sound (acoustic reflex). The intensity of the response diminishes if the stimulus is repeated (habituation). Because they are polysynaptic, extrinsic reflexes have a longer latency (stimulus-to-response interval) than intrinsic reflexes. Some important extrinsic reflexes for normal function are the postural and righting reflexes, feeding reflexes (sucking, swallowing, licking), and autonomic reflexes (p. 110).

The *flexor reflex* is triggered by noxious stimulation, e.g., from stepping on a tack. Excitatory interneurons activate spinal cord alpha-motor neurons, which, in turn, excite ipsilateral flexor muscles and simultaneously inhibit ipsilateral extensor muscles via inhibitory interneurons. Meanwhile, the contralateral extensors contract, and the contralateral flexors relax. The response does not depend on pain, which is felt only when sensory areas in the brain have been activated, by which time the motor response has already occurred. This spinal reflex arc, like that of the intrinsic muscle reflexes, is under the influence of higher motor centers.

Abnormalities of the extrinsic reflexes imply an interruption of the reflex arc or of the corticospinal tracts (which convey impulses from higher motor centers). Some clinically important extrinsic reflexes are the abdominal (T6– T12), cremasteric (L1–L2), bulbocavernosus (S3– S4), and anal wink (S3–S5) reflexes.

Reflexes that can be elicited only in the diseased state are called *pathological reflexes*. Pathological reflexes indicating dysfunction of the pyramidal (corticospinal) tract include the Babinski sign (tonic dorsiflexion of the great toe on stimulation of the lateral sole of the foot), the Gordon reflex (same response to squeezing of the calf muscles), and the Oppenheim reflex (same response to a downward stroke of the examiner's thumb on the patient's shin).

Reflexes



Motor Control

The motor system controls the timing, direction, amplitude, and force of movement through the coordinated opposing actions of agonist and antagonist muscles. It also keeps the body in a stable position through postural and righting reflexes. *Reflex movements* are involuntary, stereotyped responses to stimuli. *Rhythmic movements* have both reflex and voluntary components. *Voluntary movements* are performed at will.

Reflex Movements

Withdrawing a foot from a noxious stimulus or spreading the arms when falling are examples of reflex movements. *Intrinsic muscle reflexes* regulate muscle tone and elasticity and are important for postural control and coordination of muscle groups. Specific functions such as joint stabilization or adjustment of contraction strength are achieved with the aid of inhibitory spinal interneurons. *Extrinsic reflexes* include protective reflexes (flexor response to noxious stimulus, corneal reflex) and postural reflexes (extensor reflex, neck reflex).

Rhythmic Movements

Walking, breathing, and riding a bicycle are rhythmic movements. They are subserved both by spinal reflex arcs and by supraspinal influence from the brain stem, cerebellum, basal ganglia, and motor cortex.

Voluntary Movements

Voluntary movements depend on a sequence of contractions of numerous different muscles that is planned to achieve a desired result (motor program). Hence different parts of the body are able to carry out similar movements (motor equivalence) more or less skillfully, e.g., simultaneous rotation of the big toe, foot, lower leg, leg, pelvis and trunk. Voluntary movements incorporate elements of the basic reflex and rhythmic movement patterns: their smooth execution depends on afferent feedback from the visual, vestibular, and proprioceptive systems to motor centers in the spinal cord, brain stem, and cerebral cortex. Further modulation of voluntary movements is provided by the cerebellum and basal ganglia, whose neural output reaches

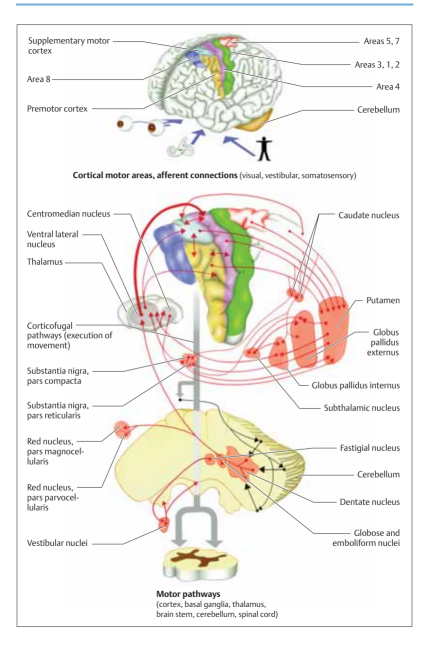
the cortex through thalamic relay nuclei. Fine motor control thus depends on the continuous interaction of multiple centers responsible for the planning (efferent copy) and execution of movement.

Motor cortex (p. 25). Voluntary movements are planned in the motor areas of the cerebral cortex. The primary motor area (area 4) regulates the force of muscle contraction and the goaloriented direction of movement; it mainly controls distal muscle groups. The supplementary motor area (medial area 6) plays an important role in complex motor planning. The premotor area (lateral area 6) receives nerve impulses from the posterior parietal cortex and is concerned with the visual and somatosensory control of movement; it mainly controls trunk and proximal limb movement.

Cerebellum (p. 54). The cerebellum coordinates limb and eye movements and plays an important role in the maintenance of balance and the regulation of muscle tone.

Basal ganglia (p. 210). The basal ganglia have a close anatomic and functional connection to the motor cortex and participate in the coordination of limb and eye movement.

Motor Control



Pyramidal Tract

Each fiber of the pyramidal tract originates in the first or upper motor neuron, whose cell body is located in the primary motor area (area 4), primary sensory areas (areas 1-3), the supplementary motor area, or the premotor area (area 6). The fibers descend through the posterior portion of the internal capsule through the cerebral peduncle, pons, and medulla, forming a small bulge (pyramid) on the anterior surface of the medulla. Most of the fibers cross the midline in the decussation of the pyramids and then descend through the spinal cord in the lateral corticospinal tract. Among the minority of fibers that do not cross in the pyramidal decussation, most continue in the ipsilateral anterior corticospinal tract, crossing the midline in the anterior spinal commissure only once they reach the level of their target motor neurons. The pyramidal tract mainly innervates distal muscle groups in the limbs. In the brain stem, the pyramidal tract gives off fibers to the motor nuclei of the cranial nerves (corticopontine and corticobulbar tracts). Fibers from the frontal eye fields (area 8) reach the nuclei subserving eye movement (cranial nerves III, IV, VI) through the pyramidal tract. The motor nuclei of cranial nerves III, IV, VI, and VII (lower two-thirds of the face) are innervated only by the contralateral cerebral cortex; thus, unilateral interruption of the pyramidal tract causes contralateral paralysis of the corresponding muscles. In contrast, the motor nuclei of cranial nerves V (portio minor), VII (frontal branch only), IX, X, XI, and XII receive bilateral cortical innervation, so that unilateral interruption of the pyramidal tract causes no paralysis of the corresponding muscles.

Nonpyramidal Motor Tracts

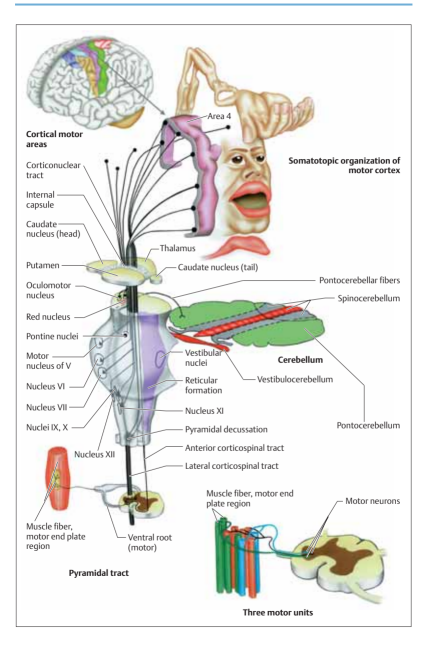
Other motor tracts lead from the cerebral cortex via the pons to the cerebellum, and from the cerebral cortex to the striatum (caudate nucleus and putamen), thalamus, substantia nigra, red nucleus, and brain stem reticular formation. These fiber pathways are adjacent to the pyramidal tract. Fibers arising from the premotor and supplementary motor areas (p.43) project ipsilaterally and contralaterally to innervate the muscles of the trunk and proximal portions of the limbs that maintain the erect body posture. Because of the bilateral innervation, paresis due to interruption of these pathways recovers more readily than distal paresis due to a pyramidal lesion. Lesions of the pyramidal tract usually involve the adjacent nonpyramidal tracts as well and cause spastic paralysis; the rare isolated pyramidal lesions cause flaccid paralysis (p. 46).

Corticopontine fibers. Corticopontine fibers originate in the frontal, temporal, parietal, and occipital cortex and descend in the internal capsule near the pyramidal tract. The pontine nuclei project to the cerebellum (p. 54).

Other functionally important tracts. The *ru-brospinal tract* originates in the red nucleus, decussates immediately, forms synapses with interneurons in the brain stem, and descends in the spinal cord to terminate in the anterior horn. Rubrospinal impulses activate flexors and inhibit extensors, as do impulses conducted in the medullary portion of the *reticulospinal tract*. On the other hand, impulses conducted in the pontine portion of the *reticulospinal tract* and in the *vestibulospinal tract* activate extensors and inhibit flexors.

Motor Unit

A motor unit is the functional unit consisting of a motor neuron and the muscle fibers innervated by it. The motor neurons are located in the brain stem (motor nuclei of cranial nerves) and spinal cord (anterior horn). The innervation ratio is the mean number of muscle fibers innervated by a single motor neuron. The action potentials arising from the cell body of a motor neuron are relayed along its axon to the neuromuscular synapses (motor end plates) of the muscle fibers. The force of muscle contraction depends on the number of motor units activated and on the frequency of action potentials. Innervation ratios vary from 3 for the extraocular muscles and 100 for the small muscles of the hand to 2000 for the gastrocnemius. The smaller the innervation ratio, the finer the gradation of force. The muscle fibers of a motor unit do not lie side by side but are distributed over a region of muscle with a cross-sectional diameter of 5-11 mm.



Paralysis Due to Upper Motor Neuron (UMN) Lesions

The clinical features of paralysis due to lesions of the pyramidal tract (upper motor neuron = UMN) depends on the anatomic site(s) of involvement of other efferent or afferent tracts and nuclei.

Impairment of fine motor function. Voluntary movement of paretic limbs requires greater effort than normal and causes greater muscular fatigue. Moreover, rapid alternating movements are slowed by hypertonia in the opposing agonist and antagonist muscles of paretic limbs. There may be *synkinesia* (involuntary movement of paretic limbs associated with other movements, e. g., yawning), undifferentiated accessory movements (*mass movements*), or *spinal automatisms* (involuntary movements triggered by somatosensory stimuli).

Paralysis. Paralysis of UMN type affects multiple (but not all) muscle groups on one side of the body. Bilaterally innervated movements (e.g., of eyes, jaw, pharynx, neck; see p. 44) may be only mildly paretic, or not at all. Paralysis that is initially total usually improves with time, but recovery may be accompanied by other motor disturbances such as tremor, hemiataxia, hemichorea, and hemiballism. Fine motor control is usually more severely impaired than strength. Neurogenic muscular atrophy does not occur in paralysis of UMN type.

Spasticity. The defining feature of spasticity is a velocity-dependent increase of muscle tone in response to passive stretch. Spasticity is usually, but not always, accompanied by hypertonia. The "clasp-knife phenomenon" (sudden slackening of muscle tone on rapid passive extension) is rare. Spasticity mainly affects the antigravity muscles (arm flexors and leg extensors).

Reflex abnormalities. The intrinsic muscle reflexes are enhanced (enlargement of reflex zones, clonus) and the extrinsic reflexes are diminished or absent. Pathological reflexes such as the Babinski reflex can be elicited.

Cerebral lesions

Monoparesis. Isolated lesions of the primary motor cortex (area 4) cause flaccid weakness of the contralateral face, hand, or leg. Lesions affecting adjacent precentral or postcentral areas,

or areas deep to the cortex, cause spasticity and possibly an associated sensory deficit. It may be difficult to determine by examination alone whether monoparesis is of upper or lower motor neuron type (p. 50). Accessory movements of antagonistic muscles are present only in paralysis of UMN type.

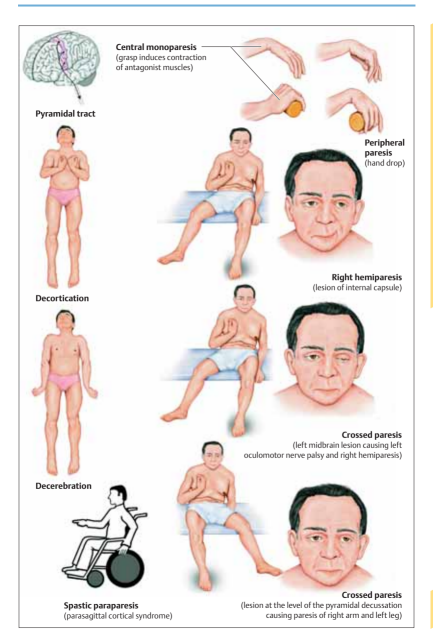
Contralateral hemiparesis. Lesions of the internal capsule cause spastic hemiparesis. Involvement of corticopontine fibers causes (central) facial paresis, and impairment of corticobulbar fibers causes dysphonia and dysphagia. Sensory disturbances are also usually present. Unilateral lesions in the rostral brain stem cause contralateral spastic hemiparesis and ipsilateral nuclear oculomotor nerve palsy (crossed paralysis). For other syndromes caused by brain stem lesions, see p. 70 ff. A rare isolated lesion of the medullary pyramid (p. 74) can cause contralateral flaccid hemiplegia without facial paralysis, or (at mid-decussational level) contralateral arm paresis and ipsilateral leg paresis (Hemiplegia alternans).

Ipsilateral paresis. Lesions of the lower medulla below the pyramidal decussation (p. 74) cause ipsilateral paralysis and spasticity (as do lesions of the lateral corticospinal tract; see p. 45).

Quadriparesis. Decortication syndrome (p. 118) is caused by extensive bilateral lesions involving both the cerebral cortex and the underlying white matter, possibly extending into the diencephalon; midbrain involvement produces the decerebration syndrome. Involvement of the pons or medulla causes an initial quadriplegia; in the later course of illness, spinal automatisms may be seen in response to noxious stimuli.

Paraparesis. In rare cases, UMN-type paralysis of both lower limbs accompanied by bladder dysfunction is caused by bilateral, paramedian, precentral cortical lesions (*parasagittal cortical syndrome*). Focal seizures may occur.

Central Paralysis



47

Spinal Cord Lesions

The site and extent of a spinal cord lesion can often be determined by clinical examination (p. 32 ff).

Paralysis. Paralysis may be of mixed upper and lower motor neuron type if the lesion affects not only the long fiber tracts but also the anterior horn cells of the spinal cord or their distal processes (root entry zone, spinal nerve roots). Reflex abnormalities are found below the level of the lesion. Central cord lesions cause both paralysis and a dissociated sensory deficit (p. 106).

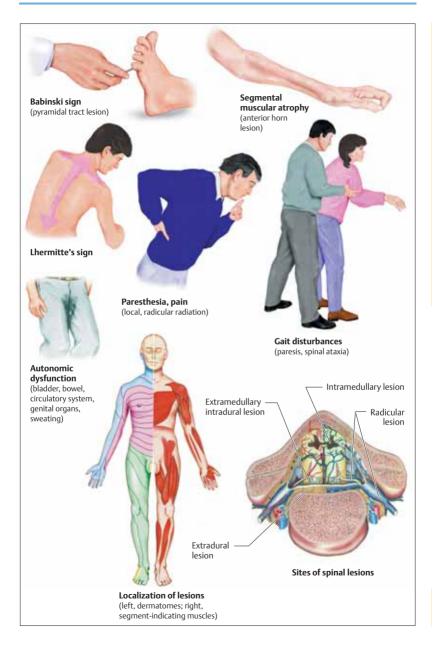
Posterior cord syndrome. Lesions of the posterior columns of the spinal cord impair vibration and position sense (p. 104 ff). Neck flexion may induce a shocklike paresthesia shooting down the back (*Lhermitte's sign*). There may be hypersensitivity to touch and noxious stimuli in areas of sensory denervation.

Autonomic dysfunction. Spinal cord transection causes acute *spinal shock*, a complete loss of autonomic function below the level of the lesion (bladder, bowel, and sexual function; vasomotor regulation; sweating). Injuries at C4 and above additionally cause respiratory paralysis. The spinal autonomic reflexes (p. 146 ff) may later recover to a variable extent, depending on the site of the lesion. Slowly progressive lesions of the caudal portion of the spinal cord (e.g., intrinsic tumor) usually come to notice because of urinary or sexual dysfunction.

Complete transection. Transection causes immediate flaccid paraplegia or quadriplegia, anesthesia and areflexia below the level of transection, bilateral Babinski signs, and spinal shock (see above). The motor and sensory impairment may begin to improve within 6 weeks if the spinal cord is incompletely transected, ultimately leading to a stable *chronic myelopathy* manifested by spastic paraparesis or guadriparesis and sensory and autonomic dysfunction. Incomplete transection. Lesions affecting only a portion of the cross-sectional area of the spinal cord cause specific clinical syndromes according to their site (pp. 32, 44, 104), of which the best known are the posterior column syndrome, the anterior horn syndrome (p. 50), the posterior horn syndrome (p. 106), the central cord syndrome (p. 106), the anterior spinal artery syndrome (p. 282), and Brown-Séguard syndrome. In the last-named syndrome, hemisection of the spinal cord causes ipsilateral spastic paresis, vasomotor paresis, anhidrosis, and loss of position and vibration sense and somatosensory two-point discrimination, associated with contralateral loss of pain and temperature sensations (the so-called dissociated sensory deficit). Cervical cord lesions. Upper cervical cord lesions at the level of the foramen magnum (p. 74) cause neck pain radiating down the arms; shoulder and arm weakness that usually begins on one side, then progresses to include the legs and finally the opposite arm and shoulder; atrophy of the intrinsic muscles of the hand; Lhermitte's sign; cranial nerve deficits (CN X, XI, XII); nystagmus; sensory disturbances on the face; and Horner syndrome. Progressive spinal cord involvement may ultimately impair respiratory function. Lesions at C1 or below do not cause cranial nerve deficits. (C1 root innervates the meninges without dermatome representation.) Lower cervical cord lesions (C5-C8) produce the signs and symptoms of complete and incomplete transection discussed above, including segmental sensory and motor deficits. If the lesion involves the spinal sympathetic pathway, Horner syndrome results.

Thoracic cord lesions. Transverse cord lesions at T1 can produce Horner syndrome and atrophy of the intrinsic muscles of the hand. Lesions at T2 and below do not affect the upper limbs. Radicular lesions produce segmental pain radiating in a band from back to front on one or both sides. Localized back pain due to spinal cord lesions is often incorrectly attributed to spinal degenerative disease until weakness and bladder dysfunction appear. Lesions of the upper thoracic cord (T1–T5) impair breathing through involvement of the intercostal muscles.

Lumbar and sacral cord injuries. Lesions at L1 to L3 cause flaccid paraplegia and bladder dysfunction (automatic bladder, p. 156). Iliopsoas weakness may make it difficult or impossible for the patient to sit. Lesions at L4 to S2 impair hip extension and flexion, knee flexion, and foot and toe movement. Lesions at S3 and below produce the conus medullaris syndrome: atonic bladder, rectal paralysis, and "saddle" anesthesia of the perianal region and inner thighs. For cauda equina syndrome, see pages 318 and 319.



Signs

Paralysis of peripheral origin can be caused by lesions of the anterior horn (lower motor neuron, LMN), nerve root, peripheral nerve, or motor end plate and must be distinguished from weakness due to disease of the muscle itself (myopathy). Apparent weakness can also be produced by tendon rupture or injury to bones and joints.

Paralysis. Paralysis is accompanied by diminution of muscle tone (flaccidity). The extent of weakness depends on the type, severity, and distribution of LMN or myopathic involvement. **Reflex abnormalities.** The intrinsic muscle reflexes are diminished or absent to a degree that may be disproportionate to the degree of weakness: in LMN-type paralysis, loss of reflexes is independent from the loss of strength; in myopathy, it parallels the weakness. Extrinsic reflexes are unaffected unless the effector muscle is atrophic. Pathological reflexes are absent.

Muscle atrophy. Muscle atrophy due to an LMN lesion may be disproportionate to the degree of weakness (either greater or less). Progressive atrophy of paralyzed muscles begins ca. 3 weeks after a peripheral nerve injury. The distribution and severity of muscle atrophy in myopathy depends on the etiology.

Spontaneous movements. Spontaneous movements are seen in affected muscles. *Fasciculations* are involuntary, nonrhythmic contractions of motor units in a relaxed muscle. They are not exclusively caused by anterior horn lesions. *Myokymia* is rhythmic contraction of muscle fibers; if the affected muscle is superficial (e.g., the orbicularis oculi), waves of muscle contraction are visible under the skin.

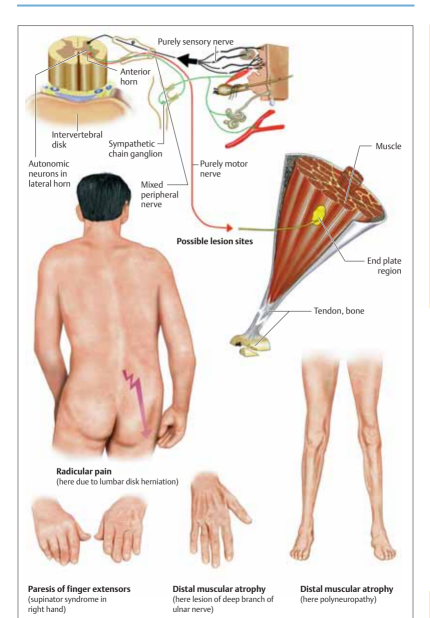
Lower Motor Neuron (LMN) Lesions

Anterior horn. Loss of motor neurons in the spinal cord paralyzes the motor units to which they belong. The flaccid segmental weakness may begin asymmetrically and is accompanied by severe muscle atrophy. There is no sensory deficit. The intrinsic reflexes of the affected muscles are lost at an early stage. The weakness may be mainly proximal (tongue, pharynx, trunk muscles) or distal (hands, calf muscles) depending on the etiology of anterior horn disease. **Radicular syndrome.** A lesion of a single ventral nerve root (caused, for example, by a herniated intervertebral disk) produces weakness in the associated myotome. Muscles supplied by multiple nerve roots are only slightly weakened, if at all, but those supplied by a single root may be frankly paralyzed and atrophic (segment-indicating muscles, cf. Table 2, p. 357). Involvement of the dorsal root produces pain and paresthesia in the associated dermatome, which may be triggered by straining (sneezing, coughing), movement (walking), or local percussion. Autonomic deficits are rare.

Peripheral nerve. Paralysis may be caused by plexus lesions (plexopathy) or by lesions of one or more peripheral nerves (mononeuropathy, polyneuropathy). Depending on the particular segment(s) of nerve(s) affected, the deficit may be purely motor, purely sensory, or mixed, with a variable degree of autonomic dysfunction.

Motor end plate. Disorders of neuromuscular transmission are typified by exercise-induced muscle fatigue and weakness. The degree of involvement of specific muscle groups (eyes, pharyngeal muscles, trunk muscles) depends on the type and severity of the underlying disease. There is no associated sensory deficit. Hyporeflexia characteristically occurs in Lambert-Eaton syndrome and is found in myasthenia gravis to an extent that parallels weakness. Autonomic dysfunction occurs in Lambert-Eaton syndrome and botulism.

Myopathy (see p. 52) and musculoskeletal lesions of the tendons, ligaments, joints, and bones may cause real or apparent weakness and thus enter into the differential diagnosis of LMN lesions. They cause no sensory deficit. Musculoskeletal lesions may restrict movement, particularly when they cause pain, sometimes to the extent that the muscle becomes atrophic from disuse. Severe autonomic dysfunction may also occur (p. 110).



Myopathy

Problems such as muscle weakness, fatigue, stiffness, cramps, tension, atrophy, pain, and involuntary movement do not necessarily signify disease of the muscle itself. Myopathy must be distinguished from neurogenic weakness of UMN or LMN type. Weakness may accompany systemic disease because of a generalized catabolic state or through a specific disease-related impairment of muscle function. Myopathy may be either primary or secondary, i.e., the product of another underlying disease. Different types of myopathy affect different muscle groups: some are generalized (congenital myopathy), while others are mainly either proximal (Duchenne type muscular dystrophy, polymyositis) or distal (myotonic dystrophy, inclusion body myositis), or mainly affect the head and face (mitochondrial myopathy). Myasthenia gravis, strictly speaking a disorder of neuromuscular transmission rather than a form of myopathy, most prominently affects the orbicularis oculi muscle: weakness increases with exercise. Muscle power is commonly graded according to the scale proposed by the British Medical Research Council (MRC) (1976):

- 0 No muscle contraction
- 1 Visible or palpable contraction, but no movement
- 2 Movement occurs, but not against gravity
- 3 Movement against gravity
- 4 Movement against gravity and additional resistance
- 5 Normal muscle power

Muscle Atrophy

Myopathy produces atrophy through the impaired development, the destruction, and the impaired regeneration of muscle fibers.

Primary (genetic) myopathies include the progressive muscular dystrophies, myotonic muscular dystrophies, congenital myopathies (e.g., central core disease, nemaline myopathy), and metabolic myopathies (Pompe disease/glycogen storage disease type II, Kearns-Sayre syndrome, carnitine deficiency).

Secondary myopathies include myositis, myopathy due to endocrine disorders (hyperthyroidism and hypothyroidism, hyperparathyroidism), and chronic toxic myopathies (alcohol, corticosteroids, chloroquine).

Disorders of Muscle Function

In these disorders, weakness is due to impaired function of the muscle fibers. Persistent weakness can lead to muscle atrophy. The episodic occurrence or worsening of muscle weakness is typical.

Primary myopathies. Hypokalemia- and hyperkalemia-related forms of paralysis belong to this group.

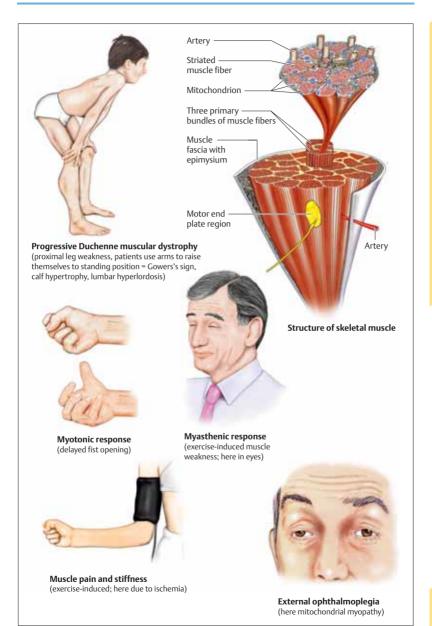
Myasthenic syndromes. Myasthenia gravis and Lambert–Eaton syndrome are characterized by abnormal fatigability of the muscles.

Postviral fatigue syndrome. Mildly increased fatigability of the muscles may persist for weeks after recovery from a viral illness.

Muscle Pain and Stiffness

Muscle pain and stiffness restrict movement, causing weakness as a secondary consequence. **Muscle pain**. Muscle pain (myalgia, p. 346) at rest and on exertion accompanies muscle trauma (muscle rupture, strain, soreness, compartment syndromes), viral myositis (influenza, Coxsackie virus, herpes simplex virus), fibromyalgia, polymyalgia rheumatica, and muscle cramps and spasms of various causes (malignant hyperthermia, carnitine palmitoyltransferase deficiency, phosphorylase deficiency/glycogen storage disease type V).

Muscle stiffness. Stiffness is prominent in congenital myotonia, neuromyotonia, and coldinduced paramyotonia.



Motor Function

The functions of the cerebellum include the control of balance, posture, gait, and goal-directed movement, and the regulation of muscle tone.

Neural Pathways

Afferent connections. The three large whitematter tracts (peduncles) of the cerebellum convey afferent input to the cerebellar cortex from the cerebral cortex (especially visual areas), pontine nuclei, the brain stem nuclei of the trigeminal, vestibular, and cochlear nerves, and the spinal cord. The superior cerebellar peduncle conveys ipsilateral proprioceptive input (p. 104) from the anterior spinocerebellar tract of the spinal cord. The middle cerebellar peduncle carries fibers of pontine origin (p. 45). The inferior cerebellar peduncle carries fibers from the vestibular nerve and nucleus to the flocculonodular lobe and fastigial nucleus, and from the contralateral inferior olive to the cerebellar hemispheres (olivocerebellar tract), as well as proprioceptive input from the posterior spinocerebellar tract (derived from muscle spindles and destined for the anterior and posterior portions of the paramedian cerebellar cortex) and fibers from the brain stem reticular formation.

Efferent connections. The cerebellar nuclei (fastigial, globose, emboliform, and dentate; p. 43) project via the (contralateral) superior cerebellar peduncle to the red nucleus, thalamus, and reticular formation. The thalamus projects in turn to the premotor and primary motor cortex, whose output travels down to the pons, which projects back to the cerebellum, forming a neuroanatomical circuit. Cerebellar output influences (ipsilateral) spinal motor neurons by way of the red nucleus and rubrospinal tract. The inferior cerebellar peduncle projects to the vestibular nuclei and brain stem reticular formation (completing the vestibulocerebellar feedback loop) and influences spinal motor neurons by way of the vestibulospinal and reticulospinal tracts.

Functional Systems

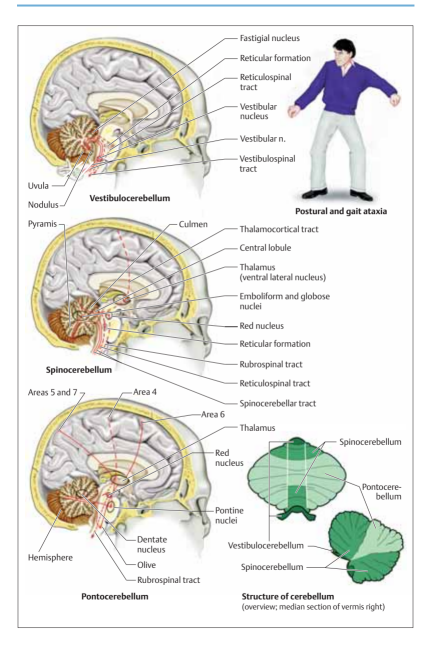
The cerebellum can be thought of as containing three separate functional components.

Vestibulocerebellum (archeocerebellum). Structures: Flocculonodular lobe and lingula. Afferent connections: From the semicircular canals and maculae (p. 56), vestibular nucleus, visual system (lateral geniculate body), superior colliculus, and striate area to the vermis. *Efferent connections*: From the fastigial nucleus to the vestibular nucleus and reticular formation. *Functions*: Control of balance, axial and proximal muscle groups, respiratory movements, and head and eye movements (stabilization of gaze). *Effects of lesions*: Loss of balance (truncal ataxia, postural ataxia > gait ataxia), nystagmus on lateral gaze, and absence of visual fixation suppression (p. 26) resulting in oscillopsia (stationary objects seem to move).

Spinocerebellum (paleocerebellum). Structures: Parts of the superior vermis (culmen, central lobule) and inferior vermis (uvula, pyramis), parts of the cerebellar hemispheres (wing of central lobule, quadrangular lobule, paraflocculus). Afferent connections: The pars intermedia receives the spinocerebellar tracts, projections from the primary motor and somatosensory cortex, and projections conveying auditory, visual, and vestibular information. Efferent connections: From the nucleus interpositus to the reticular formation, red nucleus, and ventrolateral nucleus of the thalamus, which projects in turn to area 4 of the cortex. Functions: Coordination of distal muscles, muscle tone (postural control), balance, and velocity and amplitude of saccades. Effects of lesions: Gait ataxia > postural ataxia, muscular hypotonia, dysmetria.

Pontocerebellum (neocerebellum). Structures: Most of the cerebellar hemispheres, including the declive, folium, and tuber of the vermis. Afferent connections: From sensory and motor cortical areas, premotor cortex, and parietal lobes via pontine nuclei and the inferior olive. Efferent connections: From the dentate nucleus to the red nucleus and the ventrolateral nucleus of the thalamus, and from these structures onward to motor and premotor cortex. Functions: Coordination, speed, and precision of body movement and speech. Effects of lesions: Delayed initiation and termination of movement, mistiming of agonist and antagonist contraction in movement sequences, intention tremor, limb ataxia.

Cerebellum



Labyrinth

The vestibular apparatus (labyrinth) consists of the saccule, the utricle, and three semicircular canals, each in a plane approximately at right angles to the others. The labyrinth is filled with fluid (endolymph) and has five receptor organs: the ampullary crests, which lie in a dilatation (ampulla) in front of the utricle at the end of each semicircular canal; the saccular macula (macula sacculi), a vertically oriented sensory field on the medial wall of the saccule; and the utricular macula (macula utriculi), a horizontally oriented sensory field on the floor of the utricle.

Semicircular canals. Angular acceleration is sensed by the hair cells of the ampullary crests and the gelatinous bodies (cupulae) suspended in the endolymph above them. Rotation about the axis of one of the semicircular canals causes its cupula to deflect in the opposite direction, because it is held back by the more slowly moving endolymph. With persistent rotation at a constant angular velocity (i.e., zero angular acceleration), the cupula returns to its neutral position; but if the rotation should suddenly stop, the cupula is deflected once again, this time in the direction of the original rotation, because it is carried along by the still moving endolymph. The subject feels as if he were rotating counter to the original direction of rotation and also tends to fall in the original direction of rotation

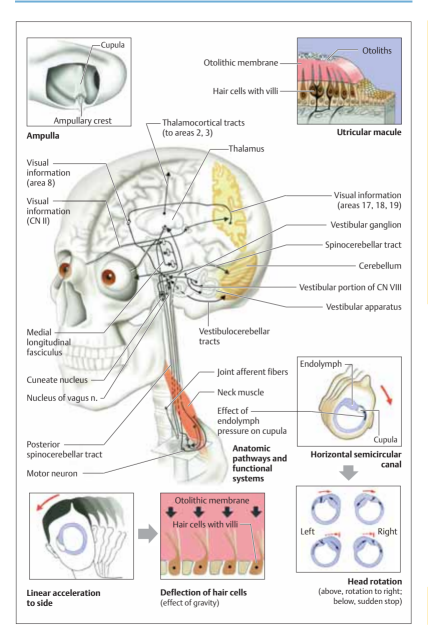
Maculae. The otolithic membrane of the saccular and utricular maculae is denser than the surrounding endolymph because of the calcite crystals (otoliths) embedded in it. *Linear acceleration* of the head thus causes relative motion of the otolithic membrane and endolymph, resulting in activation of the macular receptor cells (hair cells). The resultant forces lead to activation of the sensory receptors of the maculae.

Neural Pathways

Afferent connections. The semicircular organs project mainly to the superior and medial vestibular nuclei, the macular organs to the inferior vestibular nuclei. The vestibulocerebellum maintains both afferent and efferent connections with the vestibular nuclei; in particular, the lateral vestibular nucleus receives its major input from the paramedian region of the cerebellar cortex. Fibers reach the vestibular nucleus from the spinal cord ipsilaterally, and also bilaterally by way of the fastigial nucleus. The oculomotor nuclei project to the ipsilateral vestibular nuclei through the medial longitudinal fasciculus. The vestibular nuclei are interconnected by internuclear and commissural fibers. Efferent connections. The vestibulocerebellum projects to the ipsilateral nodulus, uvula, and anterior lobe of the vermis, and to the flocculi bilaterally. The lateral vestibulospinal tract projects ipsilaterally to the motor neurons of the spinal cord and also gives off fibers to cranial nerves X and XI. Fibers to the motor neurons of the contralateral cervical spinal cord decussate in the medial vestibulospinal tract. The medial longitudinal fasciculus (p. 86) gives off caudal fibers to the motor neurons of the cervical cord, and rostral fibers bilaterally to the nuclei subserving eye movement. Other fibers cross the midline to the contralateral thalamus, which projects in turn to cortical areas 2 and 3 (primary somatosensory area).

Functional Systems

The vestibular system provides vestibulocochlear input to the cerebellum, spinal cord, and oculomotor apparatus to enable the coordination of head, body, and eye movements. It influences extensor muscle tone and reflexes via the lateral vestibulospinal tract (postural motor system). The medial longitudinal fasciculus permits simultaneous, integrated control of neck muscle tone and eye movements. The oculomotor system (p. 86) communicates with the vestibular nuclei, the cerebellum, and the spinal cord via the medial longitudinal fasciculus and pontine projection fibers; thus the control of eye movements is coordinated with that of body movements. Proprioceptive input concerning joint position and muscle tone reaches the vestibular system from the cerebellum (p. 54). Thalamocortical connections permit spatial orientation. Phenomena such as nausea, vomiting, and sweating arise through interaction with the hypothalamus, the medullary "vomiting center," and the vagus nerve, while the emotional component of vestibular sensation (pleasure and discomfort) arises through interaction with the limbic system.



Vertigo

Patients often use the word "dizziness" nonspecifically to mean lightheadedness, unsteadiness, reeling, staggering, or a feeling of rotation. Dizziness in this broad sense has many possible causes. *Vertigo*, or dizziness in the narrow sense, is the unpleasant illusion that one is moving or that the external world is moving (so-called subjective and objective vertigo, respectively).

Pathogenesis. Vertigo arises from a mismatch between expected and received sensory input (vestibular, visual, and somatosensory) regarding spatial orientation and movement.

Cause. Vertigo occurs as a normal response to certain stimuli (physiological vertigo) or as the result of diseases (pathological vertigo) affecting the labyrinth (peripheral vestibular vertigo), central vestibular system (central vestibular vertigo), or other functional systems (nonvestibular vertigo).

Symptoms and signs. The manifestations of vertigo are the same regardless of etiology. They fall into the following categories: *autonomic* (drowsiness, yawning, pallor, sialorrhea, increased sensitivity to smell, nausea, vomiting), *mental* (decreased drive, lack of concentration, apathy, sense of impending doom), *visual* (oscillopsia = illusory movement of stationary objects), and *motor* (tendency to fall, staggering and swaying gait).

Physiological Vertigo

Healthy persons may experience vertigo when traveling by car, boat, or spaceship (*kinetosis* = *motion sickness*) or on looking down from a mountain or tall building (*height vertigo*).

Peripheral Vestibular (Labyrinthine) Vertigo (p. 88)

There is usually an acute, severe rotatory vertigo directed away from side of the labyrinthine lesion, with a tendency to fall toward the side of the lesion, horizontal nystagmus away from the side of lesion, nausea, and vomiting. Peripheral vestibular vertigo may depend on position, being triggered, for example, when the patient turns over in bed or stands up (positional vertigo), or it may be independent of position (persistent vertigo). It may also occur in attacks as episodic vertigo. Positional vertigo. Benign, paroxysmal positional vertigo (BPPV) of peripheral origin is usually due to detached otoliths of the utricular macula floating in the posterior semicircular canal (canalolithiasis). With every bodily movement, the freely floating otoliths move within the canal. under the effect of gravity. An abnormal cupular deflection results, starting 1-5 seconds after movement and lasting up to 30 seconds. The Dix-Hallpike maneuver is a provocative test for BPPV: the patient is rapidly taken from a sitting to a supine position while the head is kept turned 45° to one side. If nystagmus and vertigo ensue, they are due to canalolithiasis on the side of the ear nearer the ground. The canalith repositioning procedure (CRP), by which particles can be removed from the semicircular canal, involves repeatedly turning the patient's head to the opposite side, then back upright.

Episodic and persistent vertigo may be due to viral infection of the vestibular apparatus (vestibular neuritis, labyrinthitis) or to *Ménière disease*, which is characterized by attacks of rotatory vertigo, tinnitus, hearing loss, and ear pressure. Other causes include labyrinthine fistula and vestibular paroxysm.

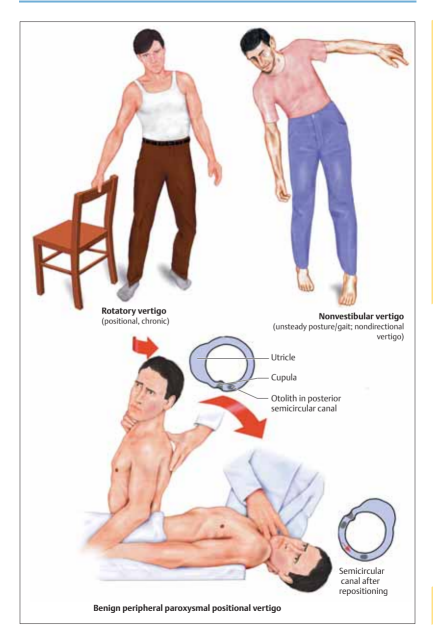
Central Vestibular Vertigo (pp. 70 ff and 88)

This type of vertigo is caused by a lesion of the vestibular nuclei, vestibulocerebellum, thalamus, or vestibular cortex, or their interconnecting fibers. Depending on the etiology (e.g., hemorrhage, ischemia, tumor, malformation, infection, multiple sclerosis, "vestibular" epilepsy, basilar migraine), vertigo may be transient or persistent, acute, episodic, or slowly progressive. It may be associated with other neurological deficits depending on the location and extent of the responsible lesion.

Nonvestibular Vertigo

Episodic or persistent nonvestibular vertigo often manifests itself as staggering, unsteady gait, and loss of balance. The possible causes include disturbances of the oculomotor apparatus, cerebellum, or spinal cord; peripheral neuropathy; intoxication; anxiety (phobic attacks of vertigo); hyperventilation; metabolic disorders; and cardiovascular disease.

Vertigo



Normal Gait

Posture. The assumption of an *upright posture* and the *maintenance of balance* (postural reflexes) are essential for walking upright. Locomotion requires the unimpaired function of the motor, visual, vestibular, and somatosensory systems. The elderly cannot stand up as quickly and tend to walk somewhat unsteadily, with stooped posture and broader steps, leading to an elevated risk of falling.

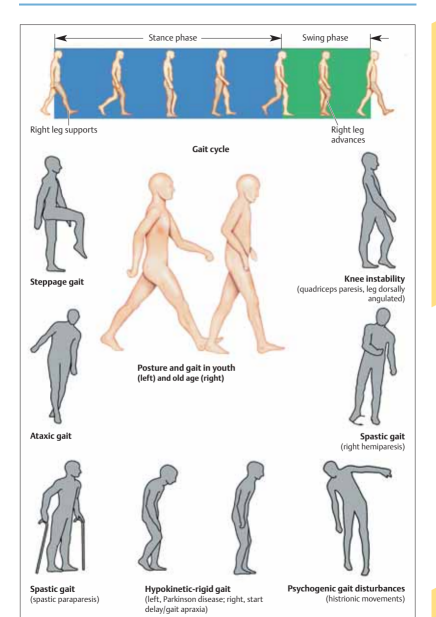
Locomotion. Normally, walking can be initiated without hesitation. The *gait cycle* (time between two successive contacts of the heel of one foot with the ground = 2 steps) is characterized by the *gait rhythm* (number of steps per unit time), the *step length* (actually the length of an entire cycle, i.e. 2 steps), and the *step width* (distance

between the lines of movement of the two heels. roughly 5-10 cm). Touchdown is with the heel of the foot. Each leg alternately functions as the supporting leg (stance phase, roughly 65% of the gait cycle), and as the advancing leg (swing phase, roughly 35% of the gait cycle). During the shifting phase, both feet are briefly in contact with the ground (double-stance phase, roughly 25% of the stance phase). Because the body's center of gravity shifts slightly to the side with each step, the upper body makes small compensatory movements to maintain balance. The arms swing alternately and opposite to the direction of leg movement. Normally, the speed of gait can be changed instantaneously. In old age, the gait sequence is less energetic and more hesitant, and turns tend to be carried out en bloc.

Gait Disturbances

Description	Related Terms	Site of Lesion	Possible Cause
Antalgic gait	Limping gait, leg differ- ence, limp	Foot, leg, pelvis, spinal column	Lumbar root lesion, bone dis- ease, peripheral nerve com- pression
Steppage gait	Foot-drop gait	Sciatic or peroneal nerve, spinal root L4/5, motor neuron	Polyneuropathy, peroneal paresis; lesions of motor neuron, sciatic nerve, or L4/5 root
Waddling gait	Duchenne gait, Trendelen- burg gait, gluteal gait	Paresis of pelvic girdle muscles (Duchenne) or of gluteal abductors (Tren- delenburg)	Myopathy, osteomalacia; le- sions of the hip joint or super- ior gluteal nerve; L5 lesion
Toe-walking		Talipes equinus, spasticity	Foot deformity, cerebral palsy, Duchenne muscular dystrophy, habit
Spastic gait	Paraspastic gait, leg cir- cumduction, spastic-ataxic gait, Wernicke–Mann gait	Pyramidal tract, extrapy- ramidal motor system (su- pratentorial, infratentorial, spinal)	Unilateral or bilateral central paralysis with spasticity, stiff- man syndrome
Ataxia of gait	Gait ataxia, staggering gait, unsteady gait, ta- betic gait, reeling gait	Peripheral nerves, posterior column of spinal cord, spinocerebellar tracts, cere- bellum, thalamus, postcen- tral cortex	Polyneuropathy, disease af- fecting posterior columns, tabes dorsalis, cerebellar le- sion, intoxication, progressive supranuclear palsy
Dystonic gait	Choreiform gait	Basal ganglia	Torsion dystonia, dopa-respon- sive dystonia, kinesiogenic paroxysmal dystonia, Hunting- ton disease
Start delay	Hypokinetic rigid gait, gait apraxia, festinating gait	Frontal lobe, basal ganglia, extensive white matter le- sions	Parkinson disease, frontal lobe lesion, normal-pressure hydro- cephalus, Binswanger disease
Psychogenic gait distur- bance	Functional gait distur- bance		Mental illness, malingering

Gait Disturbances



Tremor

Tremor, the most common movement disturbance, is an involuntary, rhythmic, oscillating movement of nearly constant amplitude. It can occur wherever movement is subserved by antagonistic muscle pairs. Different types of tremor may be classified by the circumstances in which they are activated or inhibited and by their location, frequency, and amplitude (Table 3, p. 357). Tremor amplitude is the most important determinant of disability. Parkinsonian tremor and essential tremor are the most common types.

Rest tremor occurs in the absence of voluntary movement and is aggravated by emotional stress (excitement, time pressure) and mental activity (e.g., conversing, reading a newspaper). The tremor subsides when the limbs are moved, but begins again when they return to the resting position. Rest tremor is a typical feature of parkinsonism.

Action tremors occur during voluntary movement. Postural tremor occurs during maintenance of a posture, especially when the arms are held outstretched, and disappears when the limbs are relaxed and supported. Essential tremor is a type of postural tremor. Kinetic tremor occurs during active voluntary movement; it may be worst at the beginning (initial tremor), in the middle (transitory tremor), or at the end of movement (terminal tremor). Intention tremor, the type that is worst as the movement nears its goal, is characteristic of cerebellar and brain stem lesions. Writing tremor and vocal tremor are examples of task-specific tremors. Dystonia-related tremors (e.g., in spasmodic torticollis or writer's cramp) can be suppressed by a firm grip (antagonistic maneuvers).

Frequency. The frequency of tremor in each individual case is relatively invariant and may be measured with a stopwatch or by electromyography. Different types of tremor have characteristic frequencies, listed in the table below, but there is a good deal of overlap, so that differential diagnosis cannot be based on frequency alone.

2.5–5 Hz Cerebellar tremor, Holmes tremor

- 3-6 Hz Parkinsonian tremor
- 7–9 Hz Essential tremor, postural tremor in parkinsonism
- 7–12 Hz Physiological tremor, exaggerated physiological tremor
- 12-18 Hz Orthostatic tremor

Tremor genesis. The tremor of Parkinson disease is due to rhythmic neuronal discharges in the basal ganglia (internal segment of globus pallidus, subthalamic nucleus) and thalamus (ventrolateral nucleus), which are the ultimate result of degeneration of the dopaminergic cells of the substantia nigra that project to the striatum (p. 210). Essential tremor is thought to be due to excessive oscillation in olivocerebellar circuits. which then reaches the motor cortex by way of a thalamic relay. Intention tremor is caused by lesions of the cerebellar nuclei (dentate, globose, and emboliform nuclei) or their projection fibers to the contralateral thalamus (ventrolateral nucleus, p. 54). In any variety of tremor, the abnormal oscillations are relayed from the motor cortex through the corticospinal tracts (p. 44) to the spinal anterior horn cells to produce the characteristic pattern of alternating contraction of agonist and antagonist muscles.

Tremor



Intention tremor (end tremor)

Dystonia

"Dystonia" is a general term for involuntary movement disorders involving sustained muscle contraction according to a stereotypic pattern, usually resulting in spasmodic or torsional movement and abnormal posture. Dystonic movements are usually exacerbated by voluntary activity. They may arise only during skilled activities such as writing or playing a musical instrument (action dystonia). Incomplete relief can be obtained by the avoidance of triggering activities and by the use of antagonistic maneuvers (e.g., placing the fingers on the chin, forehead or neck, or yawning, to counteract cervical dystonia). Dystonia may be classified by its distribution as focal (affects only one region of the body), segmental (two adjacent regions), multifocal (two or more nonadjacent regions), generalized, or lateralized (hemidystonia), and by its etiology as either primary (idiopathic) or secondary (symptomatic). Secondary dystonia is usually caused by a disorder of copper, lipid, or amino acid metabolism, or by a mitochondrial disorder (p. 306 ff).

Craniocervical Dystonia

Blepharospasm. Spasmodic contraction of the orbicularis oculi muscle causes excessive blinking and involuntary eye closure. It can often be accompanied by ocular foreign-body sensation and be ameliorated by distracting maneuvers, and is worse at rest or in bright light. There may be involuntary clonic eye closure, tonic narrowing of the palpebral fissure, or difficulty opening the eyes (eye-opening apraxia, p. 128). Blepharospasm may be so severe as to leave the patient no useful vision.

Oromandibular dystonia affects the perioral muscles and the muscles of mastication. In a condition named *Meige syndrome* blepharospasm is accompanied by dystonia of the tongue, larynx, pharynx, and neck.

Cervical dystonia may involve head rotation (*torticollis*), head tilt to one side (*laterocollis*), or flexion or extension of the neck (*anterocollis*, *retrocollis*), often accompanied by tonic shoulder elevation or head tremor. It may be difficult to distinguish nondystonic from dystonic head tremor; only the latter can be improved by antagonistic maneuvers. Dystonia often causes pain, usually in the neck and shoulder.

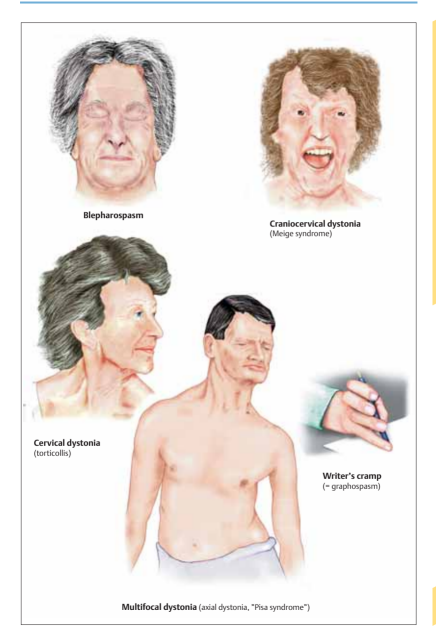
Arm and Leg Dystonia

These are most often produced by specific, usually complex, activities (*task-specific dystonia*). *Writer's cramp* (graphospasm) and musician's dystonia (for example, while playing the piano, violin, or wind instruments) are well-known examples. Toe or foot dystonia ("striatial foot") is seen in patients with Parkinson disease and dopa-responsive dystonia.

Other Types of Dystonia (see also p. 204)

In idiopathic torsion dystonia, focal dystonia of an arm or leg appears in childhood and slowly becomes generalized to include a truncal dystonia causing abnormal posture (scoliosis, kyphosis, opisthotonus). In spastic dysphonia, the voice usually sounds strained and forced, and is interrupted by constant pauses (adductor type); less commonly, it becomes breathy or whispered (abductor type). Dopa-responsive dystonia (Segawa syndrome) arises in childhood and mainly impairs gait (p. 60), to a degree that varies over the course of the day. Paroxysmal. autosomal dominant inherited forms of dystonia are characterized by recurrent dystonic attacks of variable length (seconds to hours). The attacks may be either kinesiogenic (provoked by rapid movements), in which case they usually involve choreoathetosis, or nonkinesiogenic (provoked by caffeine, alcohol, or fatigue).

Dystonia



65

Chorea

Choreiform movements are irregular, abrupt, and seemingly randomly occurring, and usually affect the distal parts of the limbs. In mild chorea, the hyperkinetic movements may be integrated in voluntary movements, such as stroking the hair. More severe cases may involve rapidly changing, bizarre body and limb postures. A combination of choreiform and (distal) dystonic movements is termed choreoathetosis. Huntington disease, an autosomal dominant disorder, is the best-known cause of chorea (p. 300, 383). Others include hereditary diseases (e.g., neuroacanthocytosis, benign hereditary chorea) and neurodegenerative diseases (e.g., Alzheimer disease, multisystem atrophy). Secondary chorea may be caused by infections (e.g., Sydenham's chorea due to streptococcal infection; herpes encephalitis, toxoplasmosis), vascular disease (e.g., lupus erythematosus, stroke), brain tumor, drug therapy (e.g., estrogen, neuroleptic drugs), or old age (senile chorea).

Hemiballism (Ballism)

Ballism consists of violent flinging movements of the limbs due to involuntary contraction of the proximal limb muscles, and usually affects only one side of the body (hemiballism). It may be continuous or occur in attacks lasting several minutes. The most common cause is an infarction or other destructive lesion of the subthalamic nucleus (STN). Diminished neural outflow from the STN leads to increased activity in the thalamocortical motor projection (p. 210).

Drug-induced Dyskinesias

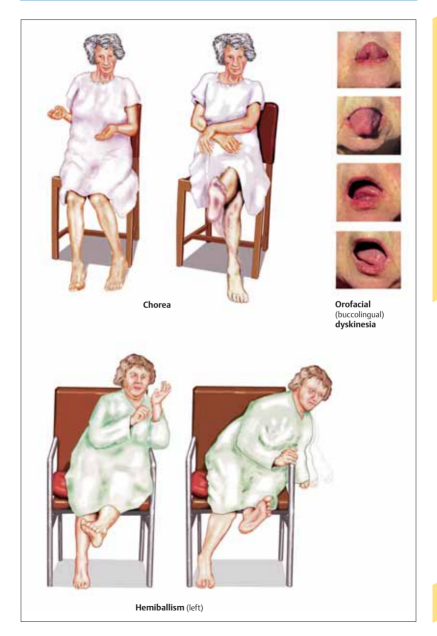
Involuntary movements of various kinds may be induced by numerous drugs, most prominently *L-dopa* and *neuroleptic drugs* including phenothiazines, butyrophenones, thioxanthenes, benzamides, and metoclopramide, all of which affect dopaminergic transmission (p. 210).

Acute dystonic reactions (p. 204) involve painful craniocervical or generalized dystonia (opisthotonus, tonic lateral bending and torsion of the trunk = "Pisa syndrome") and are treated with anticholinergic agents (e.g., biperidene).

Tardive (i.e., late) dyskinesia is a complication of long-term administration of the so-called classical neuroleptic agents. It is characterized by abnormal, stereotyped movements of the mouth, jaw, and tongue (*orofacial dyskinesia*), sometimes accompanied by respiratory disturbances, grunting, and thrusting movements of the trunk and pelvis. The same drugs may induce tardive *akathisia*, i.e., motor restlessness with a feeling of inner tension and abnormal sensations in the legs; this syndrome must be distinguished from restless legs syndrome (p. 114) and tics (p. 68). These agents also rarely induce tardive craniocervical dystonia, myoclonus, and tremor.

Myoclonus

Myoclonus consists of involuntary, brief, sudden, shocklike muscle contractions producing visible movement. It has a variety of causes and may be focal, segmental, multifocal, or generalized. Its cortical, subcortical, or spinal origin can be determined by neurophysiological testing. Attacks of myoclonus may be spontaneous or may be evoked by visual, auditory, or somatosensory stimuli (reflex myoclonus) or by voluntary movement (postural myoclonus, action myoclonus).



Physiological myoclonus. Myoclonus of variable intensity may occur normally as a person falls asleep (*sleep myoclonus*). *Hiccups* (singultus) are myoclonic movements of the diaphragm and normally cease spontaneously. (Severe, intractable hiccups, however, may be produced by lesions of the brain stem.) Normal myoclonic *startle reflexes* are to be distinguished from the rare *startle disorders* such as hyperekplexia, stiff-man syndrome, and startle epilepsy. The myoclonus that occurs in the waking phase after syncope is sometimes mistaken for an epileptic seizure.

Essential myoclonus is a rare hereditary disease characterized by persistent, very brief, multifocal myoclonic movements, accompanied by dystonia. The abnormal movements are improved by small quantities of alcohol.

Myoclonic encephalopathies. Multifocal or generalized action myoclonus is found in association with dementia and tonic-clonic seizures in various types of progressive myoclonic encephalopathy (PME), including Lafora disease, myoclonus epilepsy with ragged red fibers (MERRF syndrome), neuronal lipofuscinosis (Kufs disease) and sialidosis type I/II. Epilepsy without dementia is found in progressive myoclonus epilepsy (Unverricht-Lundborg syndrome) and progressive myoclonic ataxia.

Symptomatic myoclonus is associated with many different diseases including Alzheimer disease, corticobasal degeneration, Huntington disease, metabolic disorders (liver disease, lung disease, hypoglycemia, dialysis), encephalitis (Creutzfeldt–Jakob disease, subacute sclerosing panencephalitis), and paraneoplastic syndromes (opsoclonus-myoclonus syndrome). It can also be the result of hypoxic/ischemic brain damage (posthypoxic action myoclonus = Lance–Adams syndrome).

Asterixis consists of brief, irregular flapping movements of the outstretched arms or hands due to sudden pauses in the train of afferent impulses to muscles ("negative" myoclonus). It is not specific for any particular disease. In toxic or metabolic encephalitis, it almost always occurs together with myoclonus.

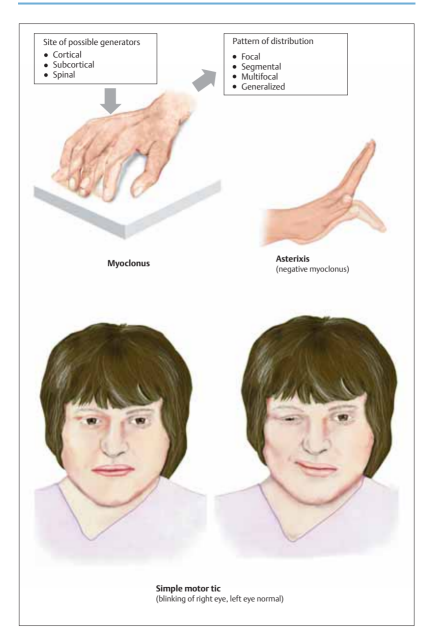
Tics

Tics are rapid, irregular, involuntary movements (motor tics) or utterances (vocal tics) that interrupt normal voluntary motor activity. They are triggered by stress, anxiety, and fatigue but may also occur at rest; they can be suppressed by a voluntary effort, but tend to re-emerge with greater intensity once the effort is relaxed. Tics are often preceded by a feeling of inner tension. They may be transient or chronic.

Simple tics. Simple motor tics involve isolated movements, e.g., blinking, twitching of abdominal muscles, or shrugging of the shoulders. Simple vocal tics may involve moaning, grunting, hissing, clicking, shouting, throat clearing, sniffing, or coughing.

Complex tics. Complex motor tics consist of stereotyped movements that may resemble voluntary movements, e.g., handshaking, scratching, kicking, touching, or mimicking another person's movements (echopraxia). Complex vocal tics may involve obscene language (coprolalia) or the repetition of another person's words or sentences (echolalia).

Gilles de la Tourette syndrome (often abbreviated to Tourette syndrome) is a chronic disease in which multiple motor and vocal tics begin in adolescence and progress over time. Other features of the disease are personality disturbances, obsessive-compulsive phenomena, and an attention deficit.



Clinical localization of brain stem lesions depends on knowledge of the tiered arrangement of cranial nerve nuclei, the intramedullary course of cranial nerve fibers, and their spatial relationship to tracts passing up and down the brain stem (see also p. 26). Lesions can be localized to the midbrain, pons, or medulla, and further classified in terms of their location in a cross-sectional plane as anterior, posterior, medial, or lateral. The "classic" brain stem syndromes are rarely seen in actual experience, as the patterns of damage tend to overlap rather than occupy discrete areas of tissue. Brain stem lesions that affect decussating neural pathways proximal to their decussation produce crossed deficits (p. 46); thus, some lesions produce ipsilateral cranial nerve palsies and contralateral hemiparesis of the limbs and trunk.

Midbrain Syndromes (Table 4, p. 358)

Lesions of the mid brain may involve its *anterior* portion (cerebral peduncle, Weber syndrome); its *medial* portion (mid brain tegmentum, Benedict syndrome), or its *dorsal* portion (midbrain tectum, Parinaud syndrome). Occlusion of the basilar artery at midbrain level causes "top of the basilar syndrome".

Pontine Syndromes

(p. 72 and Table 5, p. 359)

The syndromes produced by *anterior* and *posterior* pontine lesions are summarized in Table 5.

Paramedian Lesions

Cause. Multiple lacunar infarcts are the most common cause.

Symptoms and signs. Unilateral lesions (mediolateral or mediocentral) cause contralateral paralysis, especially in the distal limb muscles; dysarthria; and unilateral or bilateral ataxia; and, sometimes, contralateral facial and abducens palsies. *Bilateral lesions* cause pseudobulbar palsy and bilateral sensorimotor deficits.

Lateral Pontomedullary Syndrome

Cause. Infarction or hemorrhage in the territory of the posterior inferior cerebellar artery or aberrant branch of the vertebral artery.

Lesion. As in Wallenberg syndrome (p. 361) with additional involvement of the facial nerve nucleus, vestibular nerve nucleus, and inferior cerebellar peduncle.

Symptoms and signs. As in Wallenberg syndrome with additional ipsilateral findings: facial palsy (nuclear), rotatory vertigo, tinnitus, hearing loss, nystagmus, cerebellar ataxia.

Medullary Syndromes

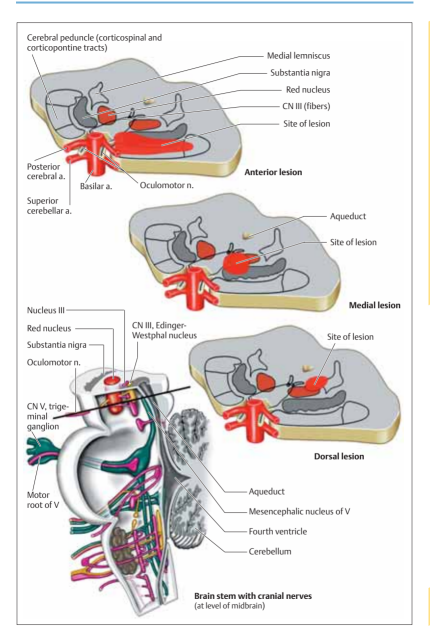
(p. 73 and Table 6, p. 361)

Lesions usually involve the medial or the lateral portion of the medulla; the lateral medullary syndrome is called Wallenberg syndrome and may be associated with various oculomotor and visual disturbances (p. 86 ff).

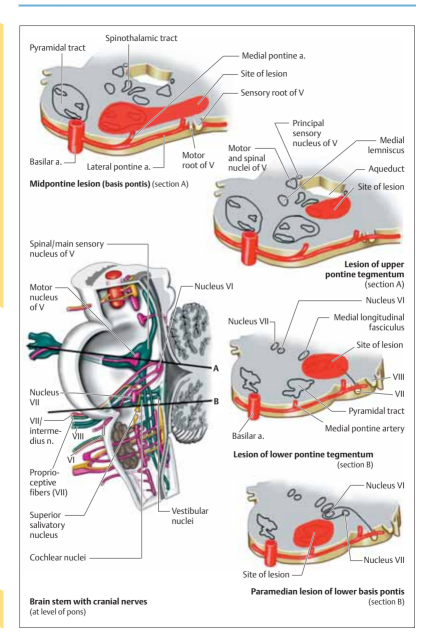
Ocular deviation. Vertical deviation (*skew deviation*) in which the ipsilateral eye is lower. Skew deviation may be accompanied by the *ocular tilt reaction*: ipsilateral head tilt, marked extorsion of the ipsilateral eye, and mild intorsion of the contralateral eye.

Nystagmus. *Positional nystagmus* may be horizontal, torsional, or mixed. *See-saw nystagmus* is characterized by intorsion and elevation of one eye and extorsion and depression of the other. **Conjugate deviation** to the side of the lesion.

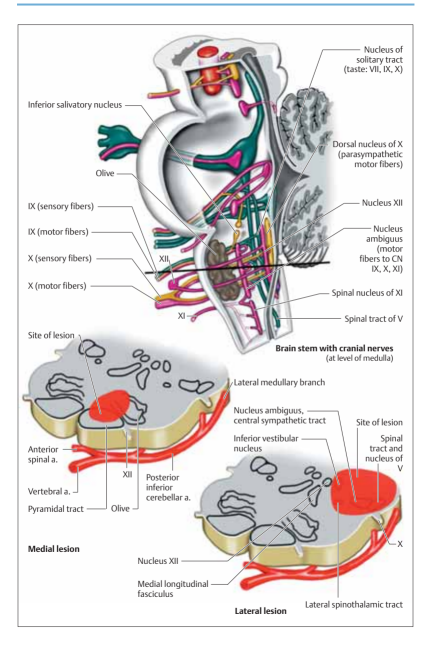
Abnormal saccades. Ocular dysmetria with overreaching (*hypermetria*) when looking to the side of lesion and underreaching (*hypometria*) when looking to the opposite side. Attempted vertical eye movements are executed with diagonal motion.



Pontine Syndromes



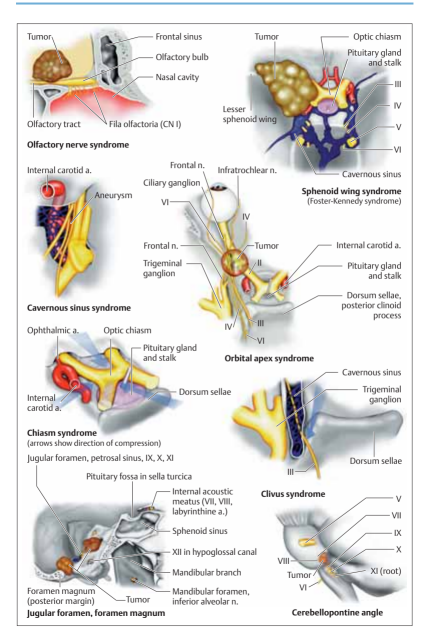
Medullary Syndromes



The site of a lesion at the base of the skull can often be deduced from the pattern of cranial nerve involvement.

Site of Lesion	Symptoms	CN ¹	Cause ²
Olfactory nerve, bulb, and tract	Anosmia, behavioral changes; may progress to Foster Ken- nedy ³ syndrome	I	Trauma, mass in anterior cranial fossa (meningioma, glioma, osteoma, abscess)
Medial sphenoid wing ⁴	Ipsilateral anosmia and optic nerve atrophy, contralateral papilledema	I, II	Medial sphenoid wing mening- ioma, mass in anterior cranial fossa
Medial/lateral sphe- noid wing ⁴	Pain in ipsilateral eye, forehead, temple; exophthalmos, diplopia	V/1, III, IV	Medial (eye pain) or lateral (tem- poral pain) sphenoid wing meningioma
Orbital apex, super- ior orbital fissure ⁵	Ipsilateral: incomplete or complete external ophthal- moplegia, sensory deficit on forehead; papilledema, visual disturbances, optic atrophy	II, III, IV, V/1, VI	Tumor (pituitary adenoma, meningioma, metastasis, naso- pharyngeal tumor, lymphoma), granuloma (TB, fungal infection, Tolosa-Hunt syndrome, arteritis), trauma, infraclinoid ICA aneurysm
Cavernous sinus ⁶	Ipsilateral symptoms and signs appear earlier than in orbital apex syndrome; exophthalmos ⁷ , Horner syndrome	III, IV, V/1, VI	Same as in orbital apex syndrome + cavernous sinus thrombosis, carotid-cavernous fistula
Optic chiasm ⁸	Visual field defects	II	See p. 80
Petrous apex ⁹	Ipsilateral facial pain (usually retro-orbital), hearing loss, sometimes also facial palsy	VI, V/1 (to V/3), VIII, (VII)	Inner ear infection, tumor, trauma
Edge of clivus ¹⁰	lpsilateral mydriasis, may pro- gress to complete oculomotor palsy	III	Intracranial hypertension (p. 158)
Cerebellopontine angle	lpsilateral hearing loss, tinnitus, deviation nystagmus, facial sensory disturbance, peripheral facial palsy/spasm, abducens palsy, ataxia, headache	VIII, V/1+2, VII, VI	Acoustic neuroma, meningioma, metastasis
Jugular foramen ¹¹	Ipsilateral: pain in region of ton- sils, root of tongue, middle ear; coughing, dysphagia, hoarse- ness, sternocleidomastoid and trapezius paresis, absence of gag reflex; sensory deficit in root of tongue, soft palate, pharynx, larynx	IX, X, XI	Metastasis, glomus tumor, trauma, jugular vein thrombosis, abscess
Foramen magnum ¹²	Same as above + ipsilateral glos- soplegia, neck pain, and local spinal symptoms (p. 48)	IX, X, XI, XII	Basilar impression, Klippel–Feil syndrome, local tumor/metastasis

1 CN = cranial nerve (unilateral CN deficits). 2 Only the most common causes are listed; other causes are possible. 3 (Foster) Kennedy syndrome (Foster is the first name). 4 Sphenoid wing syndrome. 5 Orbital apex syndrome, superior orbital fissure syndrome. 6 Cavernous sinus syndrome. 7 Patients with carotid-cavernous fistulae have pulsatile exophthalmos, conjunctival injection, and a systolic bruit that can be heard by auscultation over the eye and temple. 8 Optic chiasm syndrome. 9 Gradenigo syndrome. 10 Clivus syndrome. 11 Jugular foramen syndrome, Vernet syndrome. 12 Collet–Sicard syndrome; may be accompanied by varying degrees of dysfunction of CN IX through XII.



Olfactory epithelium. The olfactory mucosa on either side of the nasal cavity occupies an area of approximately 2.5 cm² on the roof of the superior nasal concha, extending to the nasal septum. The mucus covering the olfactory epithelium is necessary for olfactory function, because molecules interact with olfactory receptors only when they are dissolved in the mucus. Olfactory cells are bipolar sensory cells with a mean lifespan of about 4 weeks. Fine bundles of cilia project from one end of each olfactory cell into the mucus. Olfactory receptors located on the cilia are composed of specific receptor proteins that bind particular odorant molecules. Each olfactory cell produces only one type of receptor protein; the cells are thus chemotopic, i.e., each responds to only one type of olfactory stimulus. Olfactory cells are uniformly distributed throughout the olfactory mucosa of the nasal conchae.

Olfactory pathway. The unmyelinated axons of all olfactory cells converge in bundles of up to 20 fila olfactoria on each side of the nose (these bundles are the true olfactory nerves), which pass through the cribriform plate to the olfactory bulb. Hundreds of olfactory cell axons converge on the dendrites of the mitral cells of the olfactory bulb, forming the olfactory glomeruli. Other types of neurons that modulate the olfactory input (e.g., granular cells) are found among the mitral cells. Neural impulses are relayed through the projection fibers of the olfactory tract to other areas of the brain including the prepiriform cortex, limbic system, thalamus (medial nucleus), hypothalamus, and brain stem reticular formation. This complex interconnected network is responsible for the important role of smell in eating behavior, affective behavior, sexual behavior, and reflexes such as salivation. The trigeminal nerve supplies the mucous membranes of the nasal, oral, and pharyngeal cavities. Trigeminal receptor cells are also stimulated by odorant molecules, but at a higher threshold than the olfactory receptor cells.

Olfactory Disturbances (Dysosmia)

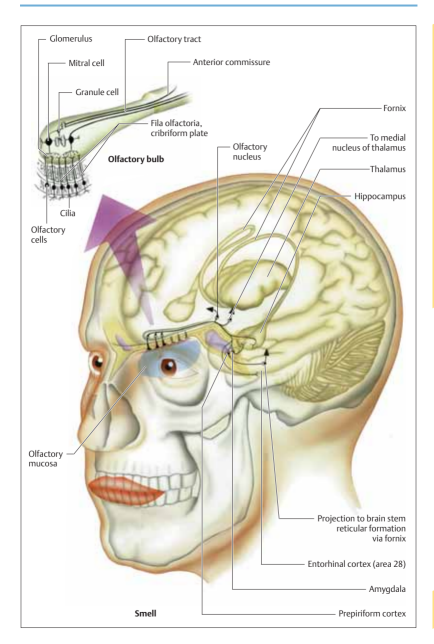
Olfactory disturbances can be classified as either *quantitative* (anosmia, hyposmia, hyperosmia) or *qualitative* (parosmia, cacosmia). Congenital olfactory disturbances manifest themselves as partial anosmia ("olfactory blindness"). The perceived intensity of a persistent odor decreases or disappears with time (olfactory adaptation). External factors such as an arid environment, cold, or cigarette smoke impair the ability to smell; diseases affecting the nasopharyngeal cavity impair both smell and taste. Odors and emotions are closely linked and can influence each other. The perception of smell may be qualitatively changed (parosmia) because of autonomic (hunger, stress) and hormonal changes (pregnancy) or disturbances such as ozena, depression, traumatic lesions, or nasopharyngeal empyema. Olfactory hallucinations can be caused by mediobasal and temporal tumors (focal epilepsy), drug or alcohol withdrawal, and psychiatric illnesses such as schizophrenia or depression.

Tests of smell. One nostril is held closed, and a bottle containing a test substance is held in front of the other. The patient is then asked to inhale and report any odor perceived. In this subjective test, odor perception per se is more important than odor recognition. Odor perception indicates that the peripheral part of the olfactory tract is intact; odor recognition indicates that the cortical portion of the olfactory pathway is also intact. More sophisticated tests may be required in some cases. Because there is bilateral innervation, unilateral lesions proximal to the anterior commissure and cortical lesions may not cause anosmia.

Anosmia/hyposmia. Unilateral anosmia may be caused by a tumor (meningioma). Korsakoff syndrome can render the patient unable to identify odors. Viral infections (influenza), heavy smoking, and toxic substances can damage the olfactory epithelium; trauma (disruption of olfactory nerves, frontal hemorrhage), tumors, meningitis, or radiotherapy may damage the olfactory pathway. Parkinson disease, multiple sclerosis, Kallmann syndrome (congenital anosmia with hypogonadism), meningoencephalocele, albinism, hepatic cirrhosis, and renal failure can also cause olfactory disturbances.

76

Smell



Taste buds. Each taste bud contains 50-150 gustatory cells. Taste buds are found on the margins and furrows of the different types of gustatory papillae (fungiform, foliate, and vallate) and are specific for one of the four primary tastes, sweet, sour, salty, and bitter. The lifespan of each gustatory cell is approximately one week. Filaments called microvilli projecting from the cells' upper poles are coated with gustatory receptor molecules. Stimulation of the gustatory cell at its receptors by the specific taste initiates a molecular transduction process. resulting in depolarization of the cell. Each taste bud responds to multiple qualities of taste, but at different sensitivity thresholds, resulting in a characteristic taste profile. For example, one papilla may be more sensitive to "sweet," another to "sour." The higher the concentration of the tasted substance, the greater the number of gustatory cells that fire action potentials. Complex tastes are encoded in the different patterns of receptor stimulation that they evoke.

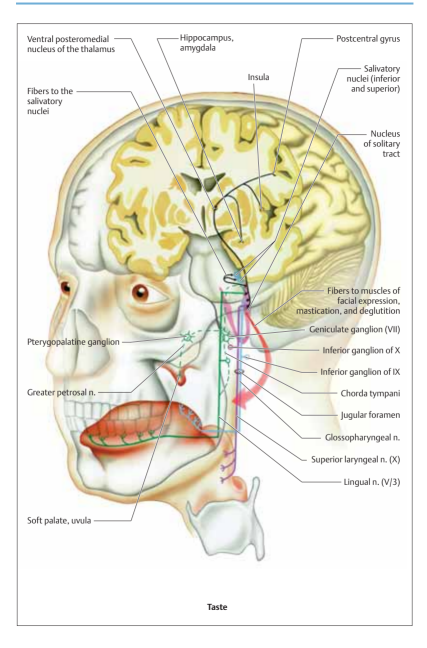
Gustatory pathway. Sensory impulses from the tongue are conveyed to the brain by three pathways: from the anterior two-thirds of the tongue via the lingual nerve (V/3) to the chorda tympani, which arises from the facial nerve (nervus intermedius); from the posterior third of the tongue via the glossopharyngeal nerve; and from the epiglottis via the vagus nerve (fibers arising from the inferior ganglion). Sensory impulses from the soft palate travel via the palatinate nerves to the pterygopalatine ganglion and onward through the greater petrosal nerve and nervus intermedius. All gustatory information arrives at the nucleus of the solitary tract, which projects, through a thalamic relay, to the postcentral gyrus. The gustatory pathway is interconnected with the olfactory pathway through the hypothalamus and amygdala. It has important interactions with the autonomic nervous system (facial sweating and flushing; salivation) and with affective centers (accounting for like and dislike of particular tastes).

Gustatory Disturbances (Dysgeusia)

When smell is impaired, the patient loses the capacity for fine differentiation of tastes but is able to distinguish the primary tastes (sweet, sour, salty, bitter). For example, chocolate pudding can be identified as "sweet" but not as "chocolate." Diminished taste (hypogeusia) is more common than complete loss of taste (ageusia).

Tests of taste. Taste thresholds on each side of the tongue are tested with the tongue outstretched. A test solution is applied to the tongue with a cotton swab for 20 to 30 seconds. The patient is then asked to point to the corresponding region of a map divided into "sweet", "sour", "salty" and "bitter" zones. The test solutions contain glucose (sweet), sodium chloride (salty), citric acid (sour), or quinine (bitter). The mouth is rinsed with water between test solutions. The taste zones are not organized in a strict topographic pattern. *Electrogustometry* can be used for precise determination of the taste thresholds but it is time-consuming and requires a high level of concentration on the part of the patient.

Ageusia/hypogeusia. Dry mouth (Sjögren syndrome), excessive alcohol consumption, smoking, spicy food, chemical burns, medications (e.g., lithium, L-dopa, aspirin, cholestyramine, amitryptiline, vincristine, carbamazepine), radiotherapy, infectious diseases (influenza), and stomatitis (thrush) can damage the taste buds. Lesions of the chorda tympani producing unilateral gustatory disturbances are seen in patients with peripheral facial palsy, chronic otitis media, and cholesteatoma. Lesions of cranial nerves V. IX. or X lead to taste distortion (especially of bitter, sour, and salty) in the posterior third of the tongue, in combination with paresthesia (sensation of burning or numbness). Gustatory disturbances may also be caused by damage to the central gustatory pathway, e.g., by trauma, brain tumors, carbon monoxide poisoning, or multiple sclerosis. The sense of taste can also change because of aging (especially sweet and sour), pregnancy, diabetes mellitus, hypothyroidism, and vitamin deficiencies (A, B₂).



79

Retina. Visible light is electromagnetic radiation at wavelengths of 400–750 nanometers. The *dioptric system* (cornea, aqueous humor of the anterior and posterior ocular chambers, pupil, lens, vitreous body) produces a miniature, upside-down mirror image of the visual field on the retina. The fovea, located in the center of the macula at the posterior pole of the eyeball, is the area of sharpest vision in daylight. Blood is supplied to the eye by the ophthalmic artery via the ciliary arteries (supplies the choroid) and the central retinal artery (supplies the retina). The optic disk, the central retinal artery that branches from it, and the central retinal vein can be examined by *ophthalmoscopy*.

Visual pathway. The visual pathway begins in the retina (first three neurons) and continues through the optic nerve to the optic chiasm, from which it continues as the optic tract to the lateral geniculate body. The optic radiation arises at the lateral geniculate body and terminates in the primary (area 17) and secondary visual areas (areas 18, 19) of the occipital lobe. The fibers of the retinal neuronal network converge at the optic disk before continuing via the optic nerve to the optic chiasm, in which the medial (nasal) fibers cross to the opposite side. The right optic tract thus contains fibers from the temporal half of the right retina and the nasal half of the left retina. The lateral geniculate body is the site of the fourth neuron of the optic pathway. Its efferent fibers form the optic radiation, which terminates in the visual cortex (striate cortex) of the occipital lobe. The central foveal area has the largest cortical representation. The visual pathway is interconnected with midbrain nuclei (medial, lateral, and dorsal terminal nuclei of the pretectal region; superior colliculus), nonvisual cortical areas (somatosensory, premotor, and auditory), the cerebellum, and the pulvinar (posterior part of thalamus).

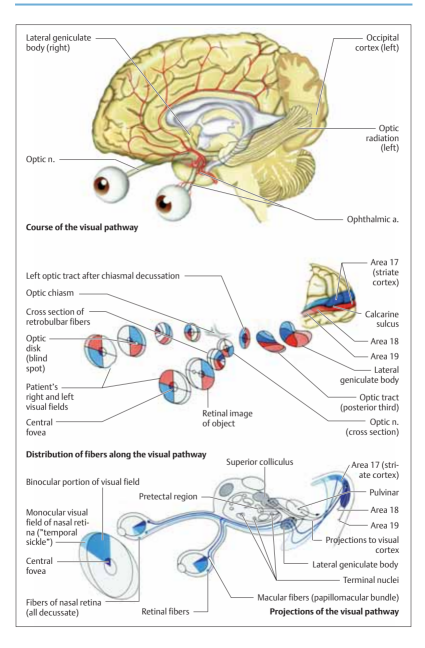
Visual field. The monocular visual field is the portion of the external world seen with one eye, and the *binocular visual field* is that seen by both eyes. The visual fields of the two eyes overlap; the overall visual field therefore consists of a central zone of clear binocular vision produced by the left and right central foveae, a peripheral binocular zone, and a monocular zone. Partial decussation at the optic chiasm brings visual information from the right (left) side of the world to the left (right) side of the brain. The visual field is topographically represented at all levels of the visual pathway from retina to cortex; lesions at any level of the pathway cause visual field defects of characteristic types. If the images on the two retinas are displaced by more than a certain threshold distance, double vision (diplopia) results. This is most commonly due to disturbances of the extraocular muscles, e. g., paralysis of one or more of these muscles (p. 86).

Stereoscopic vision. Three-dimensional visual perception (stereoscopic vision) is produced by comparison of the slightly different images in the two eyes. Stereoscopic vision is very important for depth perception, though depth can be judged to some extent, through other cues, with monocular vision alone.

Color vision. Testing of color vision requires standard definition of the colors red, blue, and green. The visual threshold for various colors. each defined as a specific mixture of the three primary colors, is determined with a standardized color perception chart. Disturbances of color vision may be due to disturbances of the dioptric system, the retina, or the visual pathway. Cortical lesions cause various kinds of visual agnosia. Lesions of area 18 may make it impossible for patients to recognize colors despite intact color vision (color agnosia), or to recognize familiar objects (object agnosia) or faces (prosopagnosia). Patients with lesions of area 19 have intact vision but cannot recognize or describe the objects that they see. Spatial orientation may be impaired (visuospatial agnosia), as may the inability to draw pictures. Persons with visual agnosia may need to touch objects to identify them.

Limbic system. Connections with the limbic system (hippocampus, amygdala, parahippocampal gyrus; p. 144) account for the ability of visual input to evoke an emotional response.

Visual pathway



Visual Field Defects

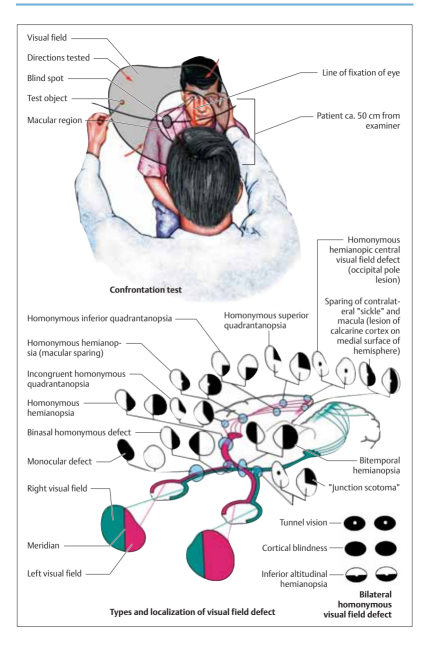
Examination. The visual fields of both eyes should always be jointly assessed. The confrontation test, in which the examiner "confronts" the patient's visual field with his or her own, intact, contralateral visual field, is used to check for visual field defects. For the test to be performed correctly, the patient and the examiner must first fixate along the same line. The examiner then slowly moves a white or red object (at least 1 cm in diameter) from the periphery of the visual field toward the center in a number of different directions, and determines where the patient can and cannot see it. Alternatively, the examiner may raise one or more fingers and ask the patient to count them (a useful test for small children, and for persons whose vision is so poor that it cannot be tested by the first method). The perceived brightness (unequal in patients with hemianopsia) of the hand in the nasal and temporal portions of the visual field is also determined. The red vision test enables the detection of a central scotoma as an area in which the red color is perceived as less intense. More detailed information can be obtained by further ophthalmological testing (Goldmann perimetry, automatic perimetry).

Visual field defect (scotoma). The thin myelinated fibers in the center of the optic nerve, which are derived from the papillomacular bundle, are usually the first to be affected by optic neuropathy (central scotoma). From the optic chiasm onward, the right and left visual fields are segregated into the left and right sides of the brain. Unilateral lesions of the retina and optic nerve cause monocular deficits, while retrochiasmatic lesions cause homonymous defects (quadrantanopsia, hemianopsia) that do not cross the vertical meridian, i.e., affect one side of the visual field only. Anterior retrochiasmatic lesions cause incongruent visual field defects, while posterior retrochiasmatic lesions lead to congruent visual field defects. Temporal lobe lesions cause mildly incongruent, contralateral, superior homonymous quadrantanopsia. Bitemporal visual field defects (heteronymous hemianopsia) have their origin in the chiasm. Unilateral retrochiasmatic lesions cause visual field defects but do not impair visual acuity. Organic visual field defects widen pregressively with the distance of test objects from the eye, whereas psychogenic ones are constant ("tubular fields"). Prechiasmatic lesions may affect the retina, papilla (= optic disk), or optic nerve. Transient

episodes of monocular blindness (amaurosis fugax) see p. 372 (table 22 a). Acute or subacute unilateral blindness may be caused by optic or retrobulbar neuritis, papilledema (intracranial mass, pseudotumor cerebri), cranial arteritis, toxic and metabolic disorders, local tumors, central retinal artery occlusion, or central retinal vein occlusion.

Chiasmatic lesions. Lesions of the optic chiasm usually produce bitemporal visual field defects. Yet, because the medial portion of the chiasm contains decussating fibers while its lateral portions contain uncrossed fibers, the type of visual field defect produced varies depending on the exact location of the lesion. As a rule, anterior chiasmatic lesions that also involve the optic nerve cause a central scotoma in the eye on the side of the lesion and a superior temporal visual field defect (junction scotoma) in the contralateral eye. Lateral chiasmatic lesions produce nasal hemianopsia of the ipsilateral eye; those that impinge on the chiasm from both sides produce binasal defects. Dorsal chiasmatic lesions produce bitemporal hemianopic paracentral scotomata. Double vision may be the chief complaint of patients with bitemporal scotomata.

Retrochiasmatic lesions. Depending on their location, retrochiasmatic lesions produce different types of homonymous unilateral scotoma: the defect may be congruent or incongruent, quadrantanopsia or hemianopsia. As a rule, temporal lesions cause contralateral superior quadrantanopsia, while parietal lesions cause contralateral inferior quadrantanopsia. Complete hemianopsia may be caused by a relatively small lesion of the optic tract or lateral geniculate body, or by a more extensive lesion more distally along the visual pathway. Sparing of the temporal sickle (p. 80) indicates that the lesion is located in the occipital interhemispheric fissure. Bilateral homonymous scotoma is caused by bilateral optic tract damage. The patient suffers from "tunnel vision" but the central visual field remains intact (sparing of macular fibers). Cortical blindness refers to subnormal visual acuity due to bilateral retrogeniculate lesions. Bilateral altitudinal homonymous hemianopsia (i.e., exclusively above or exclusively below the visual equator) is due to extensive bilateral damage to the temporal lobe (superior scotoma) or parietal lobe (inferior scotoma).



The visual axes of the eyes are directed straight ahead on primary gaze (i.e., 23° inward from the more lateral axes of the orbits). Movements of the eyes are mediated by six extraocular muscles on each side. The lateral and medial rectus muscles are responsible for horizontal eye movements. Vertical eye movements are subserved by the superior and inferior rectus as well as superior and inferior oblique muscles. The rectus muscles elevate and depress the eye when it is abducted, the oblique muscles when it is adducted. The two muscles of each synergistic pair (e.g., the left lateral rectus and right medial rectus muscles) receive equal degrees of innervation (Hering's law).

Vestibulo-ocular reflex (VOR). Impulses arising in the semicircular canals in response to rapid movement of the head induce reflex movement of the eyes in such a way as to stabilize the visual image (p. 26). For example stimulation of the horizontal semicircular canal activates the ipsilateral medial rectus and contralateral lateral rectus muscles, while inhibiting the ipsilateral lateral rectus and contralateral medial rectus muscles. The VOR makes the eyes move in the direction opposite to the head movements, at the same angular velocity.

Optokinetic reflex. Optokinetic nystagmus (OKN) is triggered by large-scale, moving visual stimuli and serves to stabilize the visual image during slow head movement. OKN is characterized by slow, gliding conjugate movement of the eyes in the direction of an object moving horizontally or vertically across the visual field, in alternation with rapid return movements in the opposite direction (saccades). OKN is intact in psychogenic (pseudo) blindness.

Fixation. Fixation is active adjustment of the gaze (with or without the aid of eye movement) to keep a visualized object in focus.

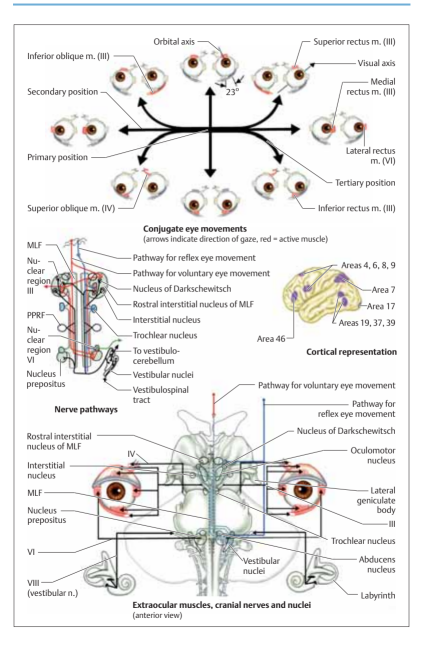
Saccades. Saccades are rapid, jerky conjugate movements of the eyes that serve to adjust or set the point of fixation of an object on the fovea. Saccades may be spontaneous, reflexive (in response to acoustic, visual, or tactile stimuli), or voluntary; the rapid phase of nystagmus is a saccade. The speed, direction, and amplitude of a saccadic movement are determined before it is carried out and cannot be influenced voluntarily during its execution. Shifts of visual fixation by more than 10° are accompanied by head move-

ments.

Slow ocular pursuit. Voluntary ocular pursuit can occur only when triggered by a moving visual stimulus (e. g., a passing car). Conversely, fixation of the gaze on a resting object while the head is moving leads to gliding eye movements. Fixation-independent ocular pursuit also occurs during somnolence and the early stages of sleep ("floating" eye movements).

Vergence movements (convergence and divergence) are mirror-image movements of the two eyes toward or away from the midline, evoked by movement of an object toward or away from the head in the sagittal plane. They serve to center the visual image on both foveae and are accompanied by an adjustment of the curvature of the lens (accommodation) to keep the object in focus.

Neural pathways. The medial longitudinal fasciculus (MLF) interconnects the nuclei of cranial nerves III, IV, and VI. The MLF also connects with fibers conveying information to and from the cervical musculature, vestibular nuclei, cerebellum, and cerebral cortex and thus mediates the coordination of eve movements with movements of the body and head. Saccades are produced by two parallel systems: Voluntary eye movements are subserved by the frontal system, which consists of the frontal eye fields (areas 4, 6, 8, 9), the supplementary eye field (area 6), the dorsolateral prefrontal cortex (area 46), and a portion of the parietal cortex (area 7). It projects to the contralateral paramedian pontine reticular formation (PPRF), which coordinates vertical and horizontal saccades. Vertical and torsional eye movements are controlled by the rostral interstitial nucleus of the MLF and by the interstitial nucleus of Cajal. Reflex eye movements are initiated in the visual cortex (area 17) and temporal lobe (areas 19, 37, 39) and modulated in the superior colliculus (collicular system). Vergence and accommodation are mediated by the pretectal area in the vicinity of the oculomotor nucleus.



Peripheral Oculomotor Disturbances

Weakness of an extraocular muscle results in diplopia, which is most pronounced in the direction of action of the affected muscle (p. 85). The cause may be a lesion in the muscle itself, in the cranial nerve that supplies it, or in the cranial nerve nucleus.

Examination. The more peripheral of the two images seen by the patient is always derived from the affected eye. The impaired eye movement may be seen directly by observation of conjugate eye movements in the nine cardinal directions of gaze (p. 85). Next, the examiner has the patient look in the direction of greatest image displacement, covers first one eye and then the other, and asks the patient each time which of the two images has disappeared. The more peripheral image disappears when the affected eve is covered. Alternatively, the patient can be asked to look at a point of light while a red glass is held in front of one eye; if the more peripheral image is red, then the eye with the glass is the affected eye. Another test is to rapidly cover and uncover one eye and then the other while the patient looks in the cardinal directions of gaze. The greatest ocular deviation and the greatest adjustment of the unaffected eye (secondary angle of deviation) occur when the patient looks in the direction of the paretic muscle. As a rule, these tests are helpful when a single muscle is acutely weak; more sophisticated ophthalmological tests are needed if the weakness is chronic or affects more than one muscle

Oculomotor nerve palsy. When a compressive lesion causes complete oculomotor nerve palsy, the patient complains of diplopia (with oblique image displacement) only when the ptotic eyelid is passively elevated. The affected eye is turned downward (action of the intact superior oblique muscle) and outward (intact lateral rectus muscle) on primary gaze, and the pupil is fixed, dilated, and irregularly shaped. The involved eye can still be abducted (intact CN VI), and looking down causes intorsion (intact CN IV). Incomplete oculomotor nerve palsy because of nuclear or myopathic lesions may differentially affect the intraocular and extraocular muscles supplied by CN III and cause different types of diplopia and pupillary disorders (p. 90). **Trochlear nerve palsy.** The affected eye points upward and toward the nose on primary gaze. Diplopia is worst when the affected eye looks toward the nose and downward.

Abducens nerve palsy. The affected eye deviates toward the nose on primary gaze. Horizontal diplopia is worst on looking toward the side of the affected eye.

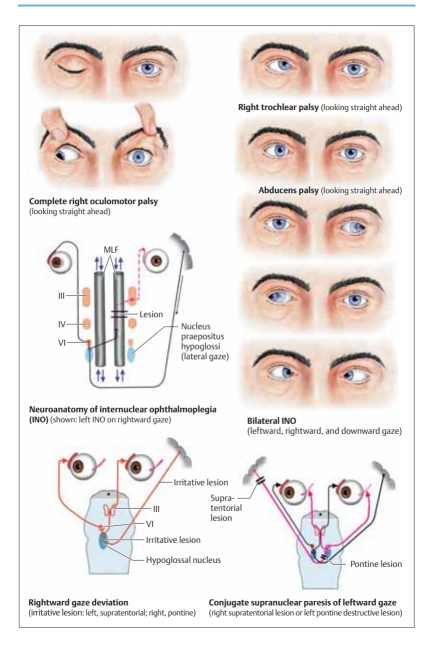
Supranuclear and Internuclear Oculomotor Disturbances

Internuclear ophthalmoplegia (INO) is characterized by inability to adduct one eye, combined with nystagmus of the other, abducted eye (dissociated nystagmus), on attempted lateral gaze. It is due to a lesion of the medial longitudinal fasciculus (MLF) on the side of the nonadducting eye and at a level between the nuclei of CN III and CN VI. Bilateral MLF lesions cause bilateral INO. Both eyes can adduct normally during convergence. More rostral lesions lead to convergence paresis without nystagmus; more caudal lesions lead to paresis of the lateral rectus muscle. Multiple sclerosis and vascular disorders are the most common causes of INO.

Unilateral pontine lesions cause ipsilateral gaze palsy (the gaze points away from the side of the lesion) but leave vertical eye movement largely intact. Co-involvement of the MLF leads to *one-and-a-half syndrome* (ipsilateral pontine gaze palsy + INO), e.g., paresis of conjugate gaze to the left and impaired adduction of the left eye on looking to the right.

Supratentorial lesions. Extensive cortical or subcortical hemispheric lesions produce *contralateral gaze palsy* (patient gazes toward the side of the lesion). Slow reflex movements of the eyes in all directions are still possible because the optokinetic reflex is not affected. In occipital lesions, the optokinetic reflex is absent; voluntary eye movements are preserved, but the eyes can no longer follow slowly moving objects. Abnormal, diffuse elevation of activity within a hemisphere (e.g., because of an epileptic seizure) causes *contralateral gaze deviation*. For further information on horizontal and verti-

For further information on horizontal and vertical gaze palsy, see page 70.



87

Nystagmus is involuntary rhythmic movement of the eyes consisting of slow movement in one direction and rapid return movement in the other. The slow component is caused by disturbances of the motor and stabilizing systems of the eye (p. 84) or because of ocular muscle paresis; the fast component represents the rapid return movement of pontine generators. Although the slow component is the actual pathological component of nystagmus, the direction of nystagmus is conventionally said to be that of its fast component, which is easier to detect. The intensity of nystagmus increases when the patient gazes in the direction of the fast component. Nystagmus can be further classified according to the type of movement as pendular, circular, or torsional (rotatory).

Examination. The examiner first observes the eyes on primary gaze, then during horizontal and vertical pursuit (fixation of gaze on a slowly moving object) and vergence. Nystagmus of labyrinthine origin is observed best with Frenzel spectacles (preventing visual fixation and giving the examiner a magnified view of the eyes). The following features of nystagmus are assessed: *positional-dependence, coordination* (conjugate, dissociated), *direction* (horizontal, vertical, rotatory, retracting, pendular), *amplitude* (fine, medium, coarse), and *frequency* (slow, moderate, fast).

Physiological Nystagmus

Physiological nystagmus serves to stabilize the visual image while the head and body are moving or when the individual looks at a moving object. The different types include *congenital nystagmus* (often X-linked recessive; fixation nystagmus is most pronounced when gazing fixedly on an object; the direction of nystagmus is usually horizontal), *spasmus nutans* (pendular nystagmus beginning in the first year of life; often accompanied by nodding of the head and torticollis; disappears spontaneously), *end-position nystagmus* (occurs during rapid movement; extreme lateral gaze; usually only a few beats), and *optokinetic nystagmus* (its absence is pathological; see p. 84).

Pathological Nystagmus

Gaze-evoked nystagmus occurs only in certain direction(s) of gaze. The main causes are drug

intoxication and brain stem or cerebellar disturbances. A slower and coarser gaze-paretic nystagmus may be seen in association with supranuclear or peripheral gaze palsy, beating in the direction of the paretic gaze. Peripheral palsy of an eye muscle may cause unilateral nystagmus of the affected eye.

Spontaneous nystagmus is that which occurs when the eyes are in the primary position; it is usually caused by vestibular dysfunction and is rarely congenital.

Peripheral vestibular nystagmus (cf. p. 58) can be seen in patients with benign paroxysmal positional vertigo, vestibular neuritis, Ménière disease, vascular compression of the vestibular nerve, and labyrinthine fistula. Nystagmus decreases on fixation and increases when fixation is blocked (lid closure, Frenzel spectacles). Most patients exhibit rotatory nystagmus that either beats continually toward the nonaffected ear, or else begins a short time after a change of position (*positional nystagmus* toward the lower ear, see p. 58).

Central vestibular nystagmus (p. 58) is caused by lesions of the brain stem (vestibular nuclei, vestibulocerebellum) or of the thalamocortical projections. It is usually accompanied by other brain stem or cerebellar signs, does not decrease on fixation, depends on the direction of gaze, and usually persists. Central positional nystagmus does not exhibit latency, is not affected by the rate of positional change, occurs with changes of position to either side, beats toward the higher ear, and is not exhaustible, stopping only when the patient is returned to the neutral position. Because positional information from vestibular, visual, and somatosensory systems is integrated in the vestibulo-ocular reflex (VOR; see pp. 26, 84), the phenomena associated with nystagmus can be explained as functional disturbances in one of the major three spatial planes of action of the VOR. Lesions cause an imbalance between the neural inputs to the VOR concerning the two sides of the affected plane. Depending on which plane is affected, the resulting nystagmus may be horizontal (horizontal plane; lesion of the vestibular nuclei), vertical (sagittal plane; pontomesencephalic, pontomedullary, or floccular lesion), or torsional (coronal plane; pontomesencephalic or pontomedullary lesion). Vertical nystagmus (upbeat or downbeat) is always due to a central lesion.

Direction of nystagmus Primary gaze Gaze-evoked nystagmus Horizontal plane = yaw (no nystagmus on primary gaze) (tendency to fall to ipsilateral side; diminished response to caloric testing in ipsilateral ear) Spontaneous nystagmus **Retraction nystagmus** (bilateral dorsal midbrain lesion) Sagittal plane = pitch Primary gaze (tendency to fall forward/backwards; "elevator" sensation) Peripheral vestibular nystagmus Frontal plane = roll (tendency to fall sideways, lateropulsion)

Central vestibular nystagmus, spatial planes

(no nystagmus on primary gaze)



Skew deviation (vertical disconjugate gaze) Vertical up- and downbeat nystagmus (brain stem lesion) The colored part of the eye, or *iris* (Greek "rainbow"), is the posterior wall of the anterior ocular chamber. Its inner edge forms the margin of the pupil. The sphincter pupillae muscle contracts the pupil, and the dilator pupillae muscle dilates it. The upper eyelid contains two muscles: the superior tarsal muscle receives sympathetic innervation, and the levator palpebrae superioris muscle is innervated by the oculomotor nerve.

Nerve Pathways

Parasympathetic fibers. The preganglionic fibers arise in the accessory oculomotor nucleus (*Edinger–Westphal nucleus*), travel in the oculomotor nerve along its outer edge, and enter the ciliary ganglion. The postganglionic fibers travel to the ciliary and sphincter pupillae muscles in the short ciliary nerves (of which there are up to 20). The parasympathetic fibers and all others on the outer aspect of CN III receive their blood supply from the pial vessels, while fibers in the interior of the nerve are supplied by the vasa nervorum.

Sympathetic fibers. The central sympathetic fibers exit from the posterolateral portion of the hypothalamus (first preganglionic neurons), then pass ipsilaterally through the tegmentum of the mid brain and pons and through the lateral medulla to form a synapse onto the second preganglionic neurons in the intermediolateral cell column of the spinal cord (ciliospinal center), at levels C8-T2. Most of the fibers exit the spinal cord with the ventral root of T1 and join with the sympathetic trunk, which lies adjacent to the pleural dome at this level. They travel with the ansa subclavia around the subclavian artery and pass through the inferior (stellate) and middle cervical ganglia to the superior cervical ganglion, where they form a (third) synapse onto the postganglionic neurons. Postganglionic fibers to the pupil travel along the course of the internal carotid artery (carotid plexus) and the ophthalmic artery, then in the nasociliary nerve (a branch of CN V) and, finally, the long ciliary nerves, which innervate the dilator pupillae muscle. Other postganglionic fibers of the sympathetic system pass to the sweat glands, the orbital muscles (bridging the inferior orbital fissure), the superior and inferior tarsal muscles, and the conjunctival vessels. Fibers to the sweat glands arise at the

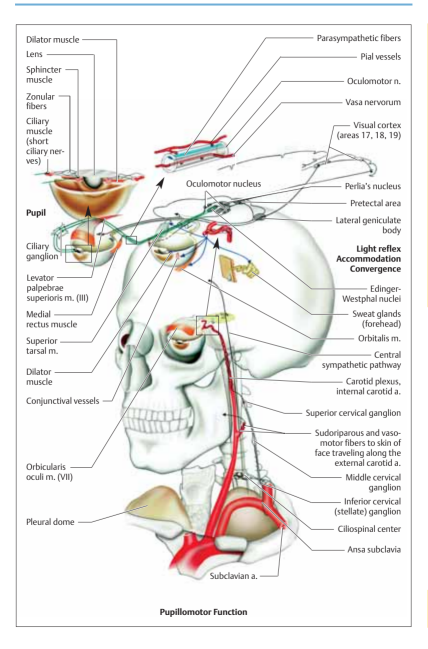
T3-T4 level and form a synapse with the third neuron in the stellate ganglion; thus, nerve root lesions at C8-T2 do not impair sweating.

Light Reflex

The light reflex regulates the diameter of the pupils according to the amount of light falling on the eye. Each pupil constricts in response to light and dilates in the dark. The afferent arm of the reflex arc consists of fibers of the optic nerve that decussate in the optic chiasm, then pass around the lateral geniculate body and terminate in the mid brain pretectal area, both ipsilaterally and contralaterally. The parasympathetic fibers are the efferent arm. The Edinger-Westphal nuclei of the two sides are connected to each other by interneurons; thus, impulses from each optic nerve arrive at both Edinger-Westphal nuclei, and light falling on one eye leads to contraction of both the ipsilateral pupil (direct light reflex) and the contralateral pupil (consensual light reflex). The pupillary diameter in moderate ambient light is normally 3-4 mm. Excessive pupillary constriction (<2 mm) is referred to as miosis, and excessive dilatation (>5 mm) as *mvdriasis*. *Anisocoria* (inequality of the diameters of the pupils) often indicates a diseased state (see below); it may be physiological but, if so, is usually mild.

The Near Response: Convergence, Pupilloconstriction, Accommodation

When a subject watches an approaching object, three things happen: the eyes converge through the action of the medial rectus muscles: the pupils constrict; and the curvature of the lens increases through the action of the ciliary muscle (accommodation). The near response may be initiated voluntarily (by squinting) but is most often the result of a reflex, whose afferent arm consists of the visual pathway to the visual cortex. The efferent arm for convergence consists of descending fibers to the pretectal convergence center (Perlia's nucleus) and onward to the oculomotor nucleus (nuclear area for the medial rectus muscles): the efferent arm for pupilloconstriction and accommodation is the parasympathetic projection of the Edinger-Westphal nucleus through the oculomotor nerve to the sphincter pupillae and ciliary muscles.



Examination. The size and shape of the pupils are first assessed in diffuse light with the patient looking at a distant object to prevent the near response. The room is then darkened and the direct light reflex of each pupil is tested at varying light intensities (by varying the distance of the lamp from the eye). If both pupils constrict when illuminated, there is no efferent pupillary defect. Next, in the swinging flashlight test, the examiner indirectly illuminates one eye with a bright light for ca. 2 seconds, then quickly switches the light to the other eye, and back again, some 5-7 times. The normal finding is that the two pupils are always of equal diameter; an abnormal finding indicates asymmetry of the afferent arm of the light reflex on the two sides, e.g., because of an optic nerve lesion (Marcus Gunn pupillary escape phenomenon). If either of these tests is abnormal, or if the pupils are significantly unequal, the near response should be tested and the direct and consensual light reflexes should be tested separately in each eye. It is easier to identify which pupil is abnormal by observing both phases of the light response (constriction and dilatation): both are slower in the abnormal pupil. In light-near dissociation, the pupils constrict as part of the near response, but not in response to light. Pharmacological pupil testing may be necessary in some cases.

Parasympathetic Denervation (Unilateral Mydriasis)

Oculomotor palsy (p. 86) is accompanied by mydriasis only when the parasympathetic fibers on the margin of the oculomotor nerve are affected. This is usually not the case in ischemic neuropathy of CN III (e.g., in diabetes mellitus), because the marginal fibers receive their blood supply from pial vessels (p. 90). A tonic pupil is a mydriatic pupil with light-near dissociation. This condition may be due to local causes (infection, temporal arteritis) or to systemic diseases such as Adie syndrome (+ reduction/absence of tendon reflexes in the legs) and Ross syndrome (+ hyporeflexia + segmental hypohidrosis). The use of anticholinergic agents (atropine eyedrops, scopolamine patch) causes iatrogenic mydriasis.

Sympathetic Denervation (Unilateral Miosis)

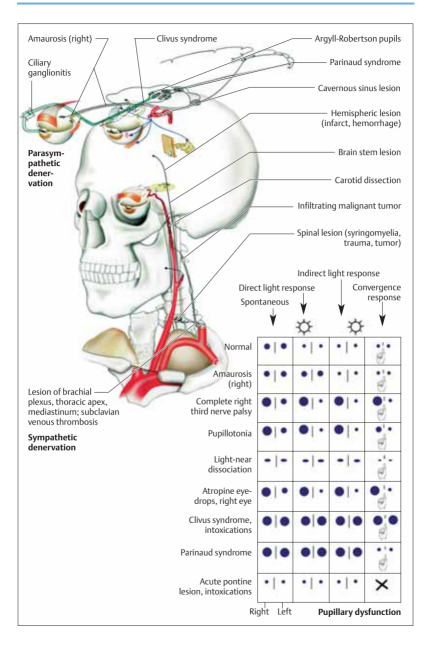
Horner syndrome is produced by a lesion at any site along the sympathetic pathway to the eve and is characterized by unilateral miosis (with sluggish dilatation) and ptosis; anhidrosis (absence of sweating) and enophthalmos are part of the syndrome but are of no practical diagnostic value. The affected pupil will fail to dilate in response to the instillation of 5% cocaine eyedrops. Preganglionic lesions (i.e., those proximal to the superior cervical ganglion) can be distinguished from postganglionic lesions by the instillation of 5% pholedrine eyedrops (at least three days after the cocaine test); the miotic pupil dilates more than the normal pupil if the lesion is preganglionic, symmetrically if it is postganglionic. Central Horner syndrome (first preganglionic neuron) may be due to lesions of hypothalamus, brain stem, or cervicothoracic spinal cord; the second preganglionic neuron may be affected by lesions of the brachial plexus, apical thorax, mediastinum, or neck: the postganglionic neuron may be affected by carotid dissection or lesions of the skull base.

Supranuclear Lesions

Lesions above the oculomotor nucleus tend to cause bilateral pupillary dysfunction; the most common cause is dorsal compression of the midbrain (Parinaud syndrome; p. 358). Neurosyphilis produces *Argyll–Robertson pupils*—unequal, irregularly miotic pupils with a variable degree of iris atrophy, and light–near dissociation.

Coma (see also p. 118)

The cause of coma may be structural, metabolic, or toxic. *Pupilloconstriction* is produced by opiates, alcohol, and barbiturates, *pupillary dilatation* by atropine poisoning (mushrooms, belladonna), tricyclic antidepressants, botulinum toxin, cocaine, and other drugs. Focal lesions (clivus, midbrain) may cause unilateral or bilateral pupillary areflexia and mydriasis. Unilateral miosis is seen in central Horner syndrome, and bilateral miosis (pinpoint pupils) in acute pontine dysfunction.



Peripheral Connections of the Trigeminal Ganglion

Ophthalmic nerve (V/1). V/1 gives off a recurrent branch to the tentorium cerebelli and falx cerebri (tentorial branch) and the lacrimal, frontal, and nasociliary nerves, which enter the orbit through the superior orbital fissure. The lacrimal nerve supplies the lacrimal gland, conjunctiva, and lateral aspect of the upper eyelid. The frontal nerve divides into the supratrochlear nerve, which supplies the inner canthus, and the supraorbital nerve, which supplies the conjunctiva, upper eyelid, skin of the forehead, and frontal sinus. Finally, the nasociliary nerve gives off branches to the skin of the medial canthus, bridge and tip of the nose, the mucous membranes of the nasal sinus (anterior ethmoid nerve) and sphenoid sinus, and the ethmoid cells (posterior ethmoid nerve).

Maxillary nerve (V/2). Before entering the foramen rotundum, V/2 gives off a *middle meningeal branch* that innervates the dura mater of the medial cranial fossa and the middle meningeal artery. Other branches innervate the skin of the zygomatic region and temple (*zygomatic nerve*), and of the cheek (*infraorbital nerve*). The infraorbital nerve enters the orbit through the inferior orbital fissure, then exits from it again through the infraorbital canal; it innervates the cheek and the maxillary teeth (superior alveolar nerve).

Mandibular nerve (V/3). V/3 gives off a meningeal branch (nervus spinosus) just distal from its exit from the foramen ovale that reenters the cranial cavity through the foramen spinosum to supply the dura mater, part of the sphenoid sinus, and the mastoid air cells. In its further course, V/3 gives off the auriculotemporal nerve (supplies the temporomandibular joint, skin of the temple in front of the ear, external auditory canal, eardrum, parotid gland, and anterior surface of the auricle), the lingual nerve (tonsils, mucous membranes of the floor of the mouth, gums of the lower front teeth, and mucosa of the anterior two-thirds of the tongue), the inferior alveolar nerve (teeth of the lower jaw and lateral gums), the mental nerve (lower lip, skin of the chin, and gums of front teeth), and the buccal nerve (buccal mucosa).

The motor root of CN V contains motor fibers from the trigeminal motor nucleus in the pons

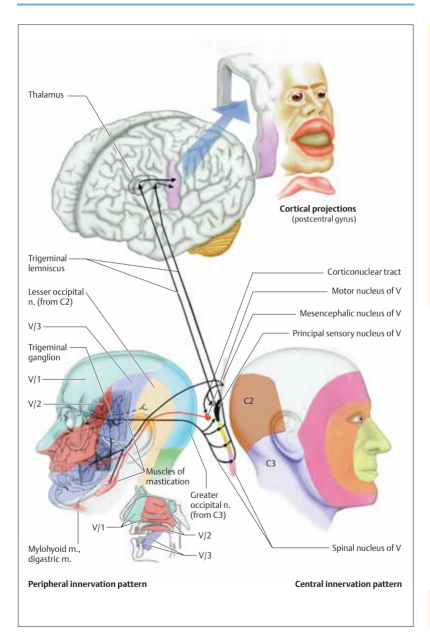
and joins the mandibular nerve to innervate the muscles of mastication (temporalis, masseter, and medial and lateral pterygoid muscles), hyoid muscles (anterior belly of the digastric muscle, mylohyoid muscle), muscles of the soft palate (tensor veli palatini muscle), and tensor tympani muscle.

Central Connections of the Trigeminal Ganglion

Sensory fibers mediating epicritic sensation terminate in the principal sensory nucleus of the trigeminal nerve, which is located in the pons. Fibers terminating in this nucleus also form the afferent arm of the corneal reflex, whose efferent arm is the facial nerve. Fibers mediating protopathic sensation terminate in the spinal nucleus of the trigeminal nerve, a column of cells that extends down the medulla to the upper cervical spinal cord. The spinal nucleus is somatotopically organized: its uppermost portion is responsible for perioral sensation, while lower portions serve progressively more peripheral areas of the face in an "onion-skin" arrangement. The caudal portion of the spinal nucleus of the trigeminal nerve also receives fibers from cranial nerves VII, IX, and X carrying nociceptive impulses from the ear, posterior third of the tongue, pharynx, and larynx.

Mesencephalic nucleus of trigeminal nerve. This midbrain nucleus, too, contains pseudounipolar neurons, whose long dendrites pass through the trigeminal ganglion without forming a synapse and carry afferent impulses from masticatory muscle spindles and pressure receptors (for regulation of the force of chewing).

Trigeminocortical tracts. Output fibers of the spinal nucleus of the trigeminal nerve decussate in the brain stem and ascend, by way of the trigeminal lemniscus (adjacent to the spinothalamic tract) and the medial lemniscus, to the ventral posteromedial (VPM) and posterior nuclei of the thalamus, where the third neuron of the sensory pathway is located. These thalamic nuclei project via the internal capsule to the postcentral gyrus. The supranuclear innervation of the motor nucleus of the trigeminal nerve is from the caudal portions of the precentral gyrus (bilaterally), by way of the corticonuclear tract.



Nerve Pathways

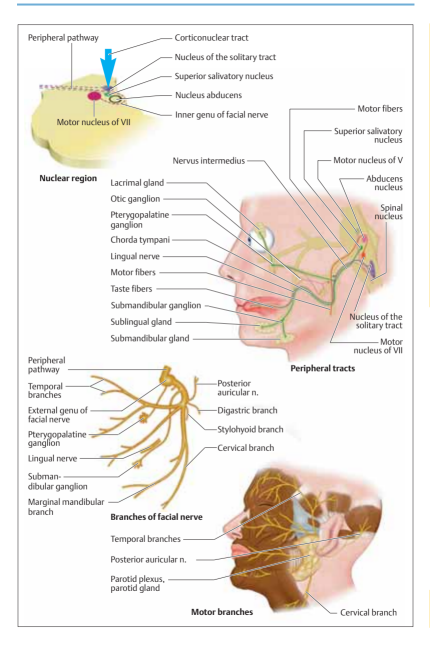
Central motor pathway. The corticonuclear tract originates in the precentral cortex (area 8), passes in front of the pyramidal tract in the genu of the internal capsule, then travels in the medial portion of the ipsilateral cerebral peduncle to reach the facial nucleus in the lower pons. The supranuclear fibers serving the upper facial muscles (frontalis and corrugator supercilii muscles, upper part of orbicularis oculi muscle, superior auricular muscle) decussate incompletely in the pons, so that these muscles have bilateral supranuclear innervation; fibers serving the remaining muscles decussate completely, so that they have contralateral innervation only. The precentral cortex is responsible for the voluntary component of facial expression, while nonpyramidal motor connections subserve the automatic and emotive components of facial expression. These anatomical facts explain the dissociated functional deficits that set supranuclear facial palsies apart from nuclear or subnuclear palsies, and enable their further differentiation into cortical and subcortical types (see below).

Peripheral motor pathway. The facial nucleus and its efferent fibers are somatotopically organized. The emerging fibers first run dorsomedially, then turn anterolaterally to pass around the abducens nucleus (inner genu of facial nerve), and exit the brain stem as the facial nerve in the cerebellopontine angle, near CN VI and VIII. The facial nerve enters the internal acoustic meatus together with the nervus intermedius and CN VIII, then leaves the meatus to enter the facial canal; it passes between the cochlea and labyrinth, then turns back again (outer genu of facial nerve). After leaving the skull at the stylomastoid foramen, it continues inside the parotid gland and gives off motor branches to all muscles of facial expression as well as the platysma, ear muscles, stapedius, digastric (posterior belly), and stylohyoid muscles. Sensory and parasympathetic fibers (nervus intermedius). Sensory fibers from the geniculate ganglion travel to the superior salivatory nucleus, nucleus of the tractus solitarius (p. 78), and spinal nucleus of the trigeminal nerve (p. 94). Taste fibers from the anterior two-thirds of the tongue (lingual nerve) and the soft palate (greater petrosal nerve) join the chorda tympani. Preganglionic parasympathetic fibers travel in the greater petrosal nerve to the pterygopalatine ganglion, from which postganglonic fibers pass to the lacrimal, nasal, and palatine glands; other preganglionic fibers travel in the chorda tympani to the submandibular ganglion, from which postganglionic fibers pass to the sublingual and submandibular glands. Connections via the contralateral medial lemniscus to the thalamus and postcentral gyrus, and to the hypothalamus, subserve reflex salivation in response to the smell and taste of food. The facial nerve carries sensory fibers from the external auditory canal, eardrum, external ear, and mastoid region (posterior auricular nerve), as well as proprioceptive fibers from the muscles it innervates.

Functional Systems

The voluntary component of facial expression is mediated by the precentral cortex, in which the face is somatotopically represented. Only the upper facial muscles have bilateral supranuclear innervation; thus, a central supranuclear facial palsy does not affect eye closure or the ability to knit one's brow. Yet facial palsy that spares the upper face is not necessarily of supranuclear origin: because the facial nucleus and nerve are also somatotopically organized, incomplete lesions of these structures may also produce a similar appearance. An important and sometimes helpful distinguishing feature is that a supranuclear palsy may affect facial expression in the lower face in a dissociated fashion. Supranuclear facial palsy due to a cortical lesion impairs voluntary facial expression, but tends to spare emotional expression (laughing, crying); that due to a subcortical lesion (e.g., in Parkinson disease or hereditary dystonia) does just the opposite.

The following reflexes are of clinical significance (A = afferent arm, E = efferent arm): orbicularis oculi reflex (blink reflex; A: V/1; E: VII); corneal reflex (A: V/1; E: VII); sucking reflex (A: V/2, V/3, XI; E: V, VII, IX, X, XII), palmomental reflex (A: thenar skin/muscles; E: VII), acoustic blink reflex (A: VIII; E: VII), visual blink reflex (A: II; E: VII), orbicularis oris reflex (snout reflex; A: V/2; E: VII).



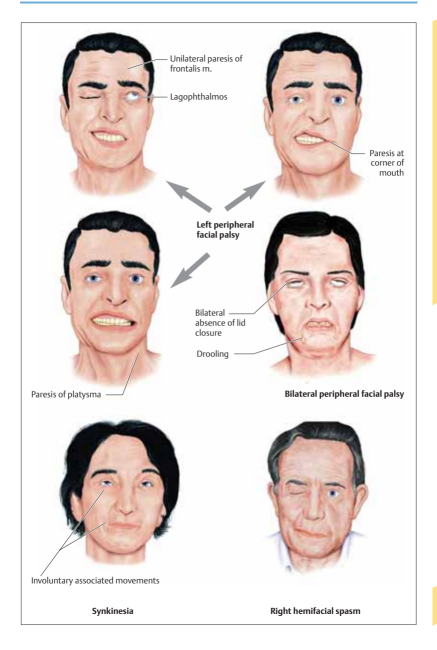
Examination. Motor function is assessed at rest (asymmetry of face/skin folds, atrophy, spontaneous movements, blink rate) and during voluntary movement (forehead, eyelids and brows, cheeks, mouth region, platysma). Trigeminal nerve dysfunction (V/1) causes unilateral or bilateral *absence* of the *blink reflex*; facial palsy may impair or abolish the blink response, but lagophthalmos persists, because the extraocular muscles are unimpaired. Similar logic applies to

other facial nerve reflexes (p. 96). If the patient complains of loss of taste, it is tested accordingly (p. 78). *Lacrimation* can be tested with the Schirmer test, which, however, is positive only if tear flow is minimal or absent. The *salivation test* is used to measure the flow of saliva from the submandibular and sublingual glands. The *stapedius reflex* is tested by measuring the contraction of the stapedius muscle in response to an acoustic stimulus.

Facial Nerve Lesions

Site of Lesion	Clinical Features	
Cortex or internal capsule	Contralateral central facial palsy (+ pyramidal tract lesion, p. 46). Emotional component of facial expression is unimpaired	
Brainstem, facial nucleus	Pontine syndrome (p. 70, 72, 359), myokymia	
Cerebellopontine angle	Ipsilateral peripheral facial palsy (+V/1–2, VI, VIII; p. 74). Hemifacial spasm ipsilateral.	
Base of skull, internal acoustic meatus	Peripheral facial palsy (+ other cranial nerve palsies; p. 74)	
Geniculate ganglion	Peripheral facial palsy, dysgeusia, hyposalivation, diminished lacrimation, ear ache, hyperacusis (due to absence of stapedius reflex)	
Facial canal distal to genicu- late ganglion	Peripheral facial palsy, dysgeusia, hyposalivation (but normal lacrimation), hyperacusis	
Proximal to stylomastoid fora- men	Peripheral facial palsy, dysgeusia, hyposalivation, intact stapedius reflex	
Stylomastoid foramen	Purely motor peripheral facial palsy	
Parotid gland, facial region	More or less complete, purely motor facial palsy; palsy due to lesions of in- dividual branches of the facial nerve	

For signs and symptoms of facial nerve lesions, see Table 7 on p. 362.



99

Perception of Sound

Sound waves enter the ear through the external acoustic meatus and travel through the ear canal to the tympanic membrane (eardrum), setting it into vibration. Vibrations in the 20-16000 Hz range (most sensitive range, 2000-5000 Hz) are transmitted to the auditory ossicles (malleus, incus, stapes). The base of the stapes vibrates against the oval window, creating waves in the perilymph in the vestibular canal (scala vestibuli) of the cochlea: these waves are then transmitted through the connecting passage at the cochlear apex (helicotrema) to the perilymph of the tympanic canal (scala tympani). (Oscillations of the round window compensate for volume changes caused by oscillations of the oval window. Sound waves can also reach the cochlea by direct conduction through the skull bone.) Migrating waves are set in motion along the basilar membrane of the cochlear duct: they travel from the stapes to the helicotrema at decreasing speed, partly because the basilar membrane is less tense as it nears the cochlear apex. These waves have their amplitude maxima at different sites along the basilar membrane. depending on frequency (tonotopicity): there results a frequency-specific excitation of the receptor cells for hearing-the hair cells of the organ of Corti, which is adjacent to the basilar membrane as it winds through the cochlea.

Cochlear Nerve

The tonotopicity of the basilar membrane causes each hair cell to be tuned to a specific sound frequency (*spectral analysis*). Each hair cell is connected to an afferent fiber of the cochlear nerve inside the organ of Corti. The cochlear nerve is formed by the central processes of the bipolar neurons of the cochlear ganglion (the first neurons of the auditory pathway); it exits from the petrous bone at the internal acoustic meatus, travels a short distance in the subarachnoid space, and enters the brain stem in the crebellopontine angle. Central auditory processing involves interpretation of the pattern and temporal sequence of the action potentials carried in the cochlear nerve.

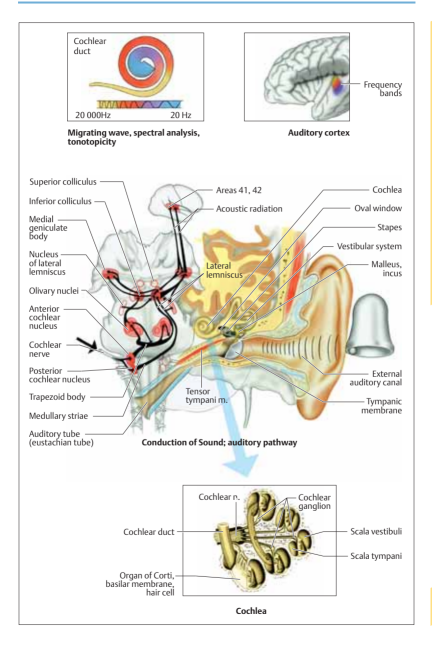
Auditory Pathway

As it ascends from the cochlea to the auditory cortex, the auditory pathway gives off collateral projections to the cerebellum, the oculomotor and facial nuclei, cervical motor neurons, and the reticular activating system, which form the afferent arm of the acoustically mediated reflexes.

Axons of the cochlear nerve originating in the cochlear apex and base terminate in the anterior and posterior cochlear nuclei, respectively. These nuclei contain the second neurons of the auditory pathway. Fibers from the posterior cochlear nucleus decussate in the floor of the fourth ventricle, then ascend to enter the lateral lemniscus and synapse in the inferior colliculus (third neuron). The inferior colliculus projects to the medial geniculate body (fourth neuron), which, in turn, projects via the acoustic radiation to the auditory cortex. The acoustic radiation passes below the thalamus and runs in the posterior limb of the internal capsule. Fibers from the anterior cochlear nucleus also decussate, mainly in the trapezoid body, and synapse onto the next (third) neuron in the olivary nucleus or the nucleus of the lateral lemniscus. This branch of the auditory pathway then continues through the lateral lemniscus to the inferior colliculus and onward through the acoustic radiation to the auditory cortex.

The primary auditory cortex (area 41: Heschl's gyrus, transverse temporal gyri) is located in the temporal operculum (i.e., the portion of the temporal lobe overlying the insula and separated from it by the sylvian cistern). Areas 42 and 22 make up the secondary auditory cortex, in which auditory signals are further processed, recognized, and compared with auditory memories. The auditory cortex of each side of the brain receives information from both ears (contralateral more than ipsilateral); unilateral lesions of the central auditory pathway or auditory cortex do not cause clinically relevant hearing loss.

100



Impairment of swallowing (deglutition) is called *dysphagia*; pain on swallowing is called *ody-nophagia*. Dysphagia or vomiting due to neuro-logical disease often causes *aspiration* (entrance of solid or liquid food into the airway below the vocal cords). *Globus hystericus* is a foreign-body sensation in the swallowing pathway independent of the act of swallowing. Despite its name, it is not always psychogenic; organic causes include Zenker diverticulum and gastroesophageal reflux.

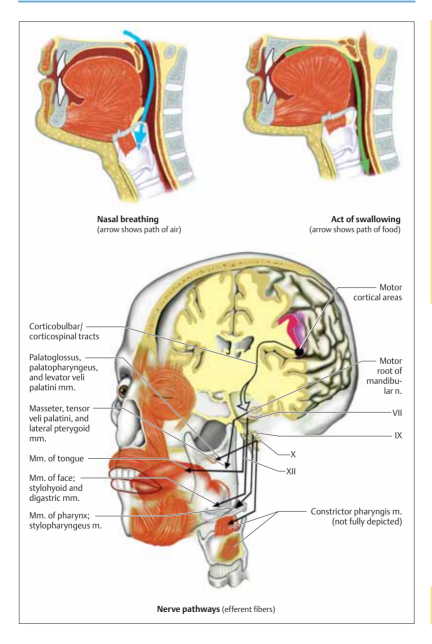
Deglutition

Mechanism. The food is ground by the teeth and moistened with saliva to form chyme, which is molded by the tongue into an easily swallowed bolus (oral preparatory phase). The tongue pushes the bolus into the oropharynx (oral phase) to initiate the reflex act of swallowing (pharyngeal phase). The lips and jaw close, the soft palate rises to seal off the nasopharynx, and the bolus bends the epiglottis backward. The bolus is pushed further back by the tongue, respiration briefly ceases, and the raised larynx occludes the airway. The upper esophageal sphincter slackens (cricopharyngeus, inferior pharyngeal constrictor, smooth muscle of upper portion of esophagus). Pressure from the tongue and pharyngeal peristalsis push the bolus past epiglottis and into the esophagus the (esophageal phase). The larynx is lowered, respiration is reinstated, and esophageal peristalsis propels the bolus into the stomach.

Nerve pathways. Fibers of CN V/2, VII, IX, and X to the nucleus ambiguus and the nucleus of the tractus solitarius (p. 78) make up the afferent arm of the swallowing reflex. The motor swallowing center (one on each side) lies adjacent to these nuclei and is associated with the upper medullary reticular formation: it coordinates the actions of the numerous muscles involved in swallowing. Efferent signals reach these muscles through CN V/3, VII, IX, X, and XII. Crossed and uncrossed supranuclear innervation is derived from the cerebral cortex (precentral and postcentral gyri, frontoparietal operculum, premotor cortex, and anterior insular region). Spinal motor neurons also participate (C1-C4).

Neurological Disturbances of Deglutition (See Table 8 on p. 362)

The disturbance usually manifests itself at the beginning of the act of swallowing (e.g., a feeling of food stuck in the throat, the escape of liquid or solid food through the nose, choking, coughing). Associated inflammation of the swallowing pathway may cause odynophagia. Chronic dysphagia causes inadequate nutrition and weight loss. Neurogenic dysphagia usually impairs the swallowing of liquids more than solids; soft, chilled foods (like pudding or vogurt) are often easier to swallow. Sensory disturbances in the larynx and trachea, a diminished cough reflex, and muscle weakness may cause aspiration, sometimes unremarked by the patient (silent aspiration). The diagnostic evaluation of dysphagia may require special tests such as radiocinematography, video endoscopy, manometry, and pH measurement.



There are two functionally and anatomically distinct types of somatic sensation and pain. The spatially and temporally precise perception of light tactile, noxious, and temperature stimuli is called *epicritic sensation*, and the more diffuse perception of stronger tactile, noxious, and temperature stimuli is called *protopathic sensation*. Sensation in the deep tissues (muscles, viscera) is predominantly protopathic.

Receptors

Sensory stimuli affect the nervous system by physically interacting with receptors. Exteroceptors respond to external stimuli (mechanical, thermal, optic, acoustic, olfactory, gustatory); interoceptors respond to internal stimuli (stretch, pressure, chemical irritation of internal organs). A stimulus activates a receptor only if it is sufficiently intense (above threshold). Receptors are classified according to their activating stimuli: mechanoreceptors (pressure, touch; proprioceptive sensations such as joint postion, muscle contraction, muscle stretch; hearing, sense of balance), thermoreceptors (heat, cold), chemoreceptors (pain, smell, itch, taste), and photoreceptors (light). Cutaneous receptors include both "free" nerve endings and specially adapted receptors (e.g., corpuscles of Meissner and Vater-Pacini). The former type mainly subserve pain and temperature sense, the latter tactile sensation (touch, pressure, vibration). In hair-covered skin there are tactile receptors around the hair roots.

Nerve Pathways

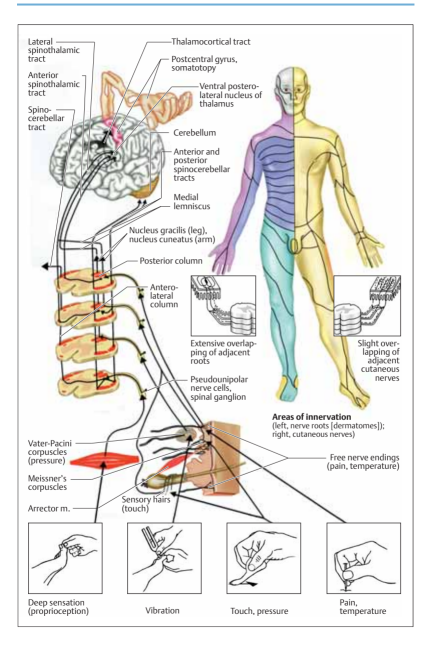
From the receptor, information is transmitted to the afferent fibers of the pseudounipolar spinal ganglion cells, whose efferent fibers reach the spinal cord by way of the dorsal root. A synapse onto a second neuron in the sensory pathway is made either immediately, in the posterior horn of the spinal cord (protopathic system), or more rostrally, in the brain stem (epicritic/lemniscal system). The highest level of the somatosensory pathway is the contralateral primary somatosensory cortex. The somatotopic organization of the somatosensory pathway is preserved at all levels. Fibers mediating sensation in the legs are in the fasciculus gracilis (medial), while those for the arms are in the fasciculus cuneatus (lateral). These fibers synapse onto the second sensory neuron in the corresponding somatosensory nuclei of the lower medulla (nucleus gracilis, nucleus cuneatus), which emit fibers that decussate and ascend in the contralateral medial lemniscus to the thalamus (ventral posterolateral nucleus, VPL). VPL projects to the postcentral gyrus by way of the internal capsule.

Anterolateral column (protopathic system). Fibers of the protopathic pathway for somatic sensation (strong pressure, coarse touch) enter the spinal cord through the dorsal root and then ascend two or more segments before making a synapse in the ipsilateral posterior horn. Fibers originating in the posterior horn decussate in the anterior commissure of the spinal cord and enter the anterior spinothalamic tract, which is somatotopically arranged: fibers for the legs are anterolateral, fibers for the arms are posteromedial. The anterior spinothalamic tract traverses the brain stem adjacent to the medial lemniscus and terminates in VPL, which, in turn, projects to the postcentral gyrus. The protopathic pathway for pain (as well as tickle, itch, and temperature sensation) is organized in similar fashion: Central fibers of the first sensory neuron ascend 1 or 2 segments before making a synapse in the substantia gelatinosa of the posterior horn. Fibers from the posterior horn decussate and enter the lateral spinothalamic tract, which, like the anterior spinothalamic tract, projects to VPL; VPL projects in turn to the postcentral gyrus.

Spinocerebellar tracts (spinocerebellar system). These tracts mediate proprioception. Fibers originating from muscles spindles and tendon organs make synapses onto the neurons of *Clarke's column* within the posterior horn at levels T1–L2, whose axons form the posterior spinocerebellar tract (ipsilateral) and the anterior spinocerebellar tract (both ipsilateral and contralateral). These tracts terminate in the spinocerebellum (p. 54).

Posterior column (epicritic/lemniscal system).

Sensation



105

Examination. Somatic sensation is tested with the patient's eyes closed. The examiner tests each primary modality of *superficial sensation* (touch, pain, temperature), the patient's ability to distinguish different qualities of each modality (sharp/blunt, hot/cold, different intensities, two-point discrimination), and more complex sensory modalities (stereognosis, graphesthesia). Next, sensation to pressure and vibration stimuli are tested, as is acrognosis (posture sense), to evaluate *proprioception*. Sensory disturbances commonly cause disturbances of posture (tests: Romberg test, standing on one leg) or gait (p. 60).

Interpretation of findings. There is a wide range of normal findings. Apparent abnormalities should be interpreted in conjunction with findings of other types, such as abnormal reflexes or paresis. Sensory dysfunction may involve not only a diminution or absence of sensation (*hypesthesia, anesthesia*), but also sensations of abnormal type (*paresthesia*, such as prickling or formication) or spontaneous pain (*dysesthesia*, often of burning type). Patients often use the colloquial term "numbness" to mean hypesthesia, anesthesia, or paresthesia; the physician should ask specific questions to determine what is meant.

Clinical Features Site of Lesion Possible Causes¹ Localized sensory disturbance (not in a Cutaneous nerves/ Skin lesions, scars, lepromatous leprosy (dissociated sensory deficit³ distally in the dermatomal or peripheral nerve distribureceptors tion)2 limbs, tip of nose, external ear) Often pain and paresthesia at first, then Distal peripheral Mononeuropathy (compression, tumor), sensory deficit, in a distribution dependnerve mononeuritis multiplex (involvement of ing on the site of the lesion multiple peripheral nerves by vasculitis, diabetes mellitus, etc.) Distal symmetrical sensory disturbances Distal peripheral Polyneuropathy (diabetes mellitus, alconerves hol, drug/toxic, Guillain-Barré syndrome) Bilateral symmetrical or asymmetrical Peripheral nerves. Diabetes mellitus thigh pain lumbar plexus Multiple sensory and motor deficits in a Plexus Trauma, compression, infection, ischemia, sinale limb tumor, metabolic disturbance Unilateral or bilateral, monoradicular or Nerve root Herniated disk, herpes zoster, Guillainpolyradicular deficits Barré syndrome, tumor, carcinomatous meningitis, paraneoplastic syndrome Spinal ataxia, incomplete or complete Spinal cord Vascular, tumor, inflammatory/multiple cord transection syndrome (p. 48) sclerosis, hereditary, metabolic disease, trauma, malformation Loss of position and vibration sense in the Craniocervical junc-Tumor, basilar impression upper limbs and trunk, Lhermitte's sign tion Contralateral dissociated or crossed Brainstem Vascular, tumor, multiple sclerosis sensory deficit (p. 70 ff) Contralateral paresthesia and sensory Thalamus Vascular (p. 170), tumor, multiple sclerodeficits, pain, loss of vibration sense Paresthesia, contralateral sensory deficits Postcentral cortex Vascular, tumor, trauma (astereognosis, loss of position sense and

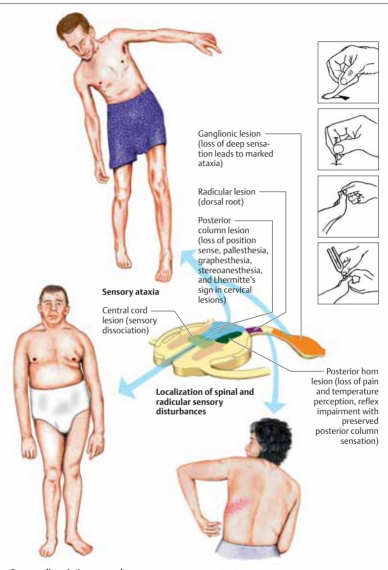
Localization of Sensory Disturbances

two-point discrimination, inability to localize a stimulus, agraphesthesia)

1 The listing of possible causes is necessarily incomplete. 2 May be factitious or psychogenic.

3 Impairment or loss of pain and temperature sensation with preserved touch sensation.

Sensory Disturbances



Sensory dissociation, muscular atrophy, scoliosis due to syringomyelia

Radicular sensory disturbances and pain in herpes zoster Sensation

Pain is an unpleasant sensory and emotional experience associated with actual or potential tissue damage, or described in terms of such damage (International Association for the Study of Pain).

Pathogenesis

Pain results from the interaction of a noxious (i.e., pain-producing) stimulus with a receptor, and the subsequent transmission and processing of pain-related signals in the PNS and CNS; the entire process is called *nociception*. Pain evokes a behavioral response involving nocifensor activity as well as motor and autonomic reflexes.

Pain reception. Nociceptors for mechanical, thermal, and chemical stimuli are found in all body organs except the brain and spinal cord. By releasing neuropeptides, the nociceptors can produce a neurogenic sterile inflammatory response that enhances nociception (*peripheral sensitization*).

Pain transmission. Nociceptive impulses travel in peripheral nerves to the posterior horn of the spinal cord. Here, the incoming information is processed by both pain-specific and nonspecific (wide dynamic range) neurons. Central sensitization processes arising at this level may lower the nociceptor threshold and promote the development of chronic pain (such as phantom limb pain after amputation). Ascending impulses reach the brain through the spinothalamic and spinoreticular tracts as well as other pathways to a number of different brain regions involved in nociception.

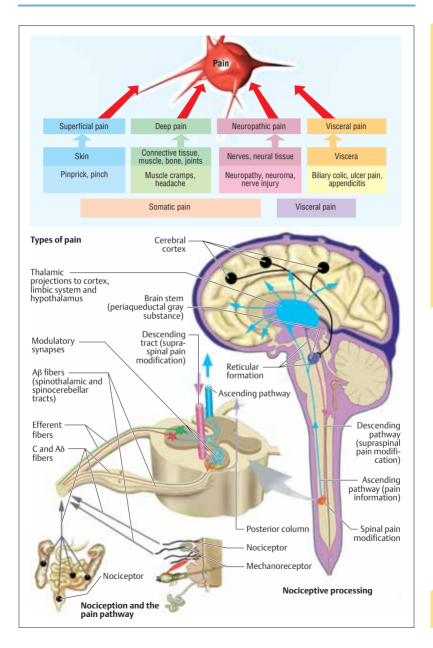
Pain processing. The reticular formation regulates arousal reactions, autonomic reflexes, and emotional responses to pain. The *thalamus* relays and differentiates nociceptive stimuli. The *hypothalamus* mediates autonomic and neuroendocrine responses. The *limbic system* (p. 144) mediates emotional and motivation-related aspects of nociception. The *somatosensory cortex* is mainly responsible for pain differentiation and localization. *Descending pathways* that originate in these CNS areas also modulate nociception.

Neurotransmitters and neuropeptides are involved in nociception on different levels. Various neurotransmitters and neuropeptide systems play a role in the mechanism of action of one or more currently used analgesic agents (effective drugs in parantheses): glutamate (memantine); substance P (capsaicin); histamine (antihistamines); serotonin/norepinephrine (antidepressants); GABA (baclofen, diazepam); prostaglandins (nonsteroidal antiinflammatory drugs); enkephalin, endorphin, dynorphin (opiates, opioids).

Types of Pain (See Table 9, p. 363)

Nociceptive pain, the "normal" type of pain, is that which arises from actual or potential tissue damage and results from the activation of nociceptors and subsequent processing in an intact nervous system. Somatic pain is the variety of nociceptive pain mediated by somatosensory afferent fibers; it is usually easily localizable and of sharp, aching, or throbbing quality. Postoperative, traumatic, and local inflammatory pain are often of this variety. Visceral pain is harder to localize (e.g., headache in meningitis, biliary colic, gastritis, mesenteric infarction) and may be dull, cramplike, piercing, or waxing and waning. It is mediated peripherally by C fibers and centrally by spinal cord pathways terminating mainly in the limbic system. This may explain the unpleasant and emotionally distressing nature of visceral pain. Visceral pain may be felt in its site of origin or may be referred to another site (e.g., from the diaphragm to the shoulder).

Neuropathic pain is that which is caused by damage to nerve tissue. It is always referred to the sensory distribution of the affected neural structure: e.g., calf pain in S1 radiculopathy, frontal headache in tentorial meningioma, unilateral bandlike abdominal pain in schwannoma of a thoracic spinal nerve root. (Note that neuropathic pain is not necessarily due to neuropathy. The less misleading synonym "neurogenic pain" is not as widely used.)



Head's Zones

Visceral pain is not felt in the internal organ where it originates, but is rather referred to a cutaneous zone (of Head) specific to that organ. This phenomenon is explained by the arrival of sensory impulses from both the internal organ and its related zone of Head at the posterior horn at the same level of the spinal cord; the brain thus (mis)interprets the visceral pain as originating in the related cutaneous zone. The pain may be described as burning, pulling, pressure, or soreness, and there may be cutaneous hyperesthesia to light touch. Certain etiologies (e.g., angina pectoris, cholecystitis, gastric ulcer, intestinal disease) can produce ipsilateral mydriasis. In addition to the zones of Head, referred pain may also be felt in muscles and connective tissue (pressure points, as in Blumberg's sign or McBurney's point). Physicians should beware of mistaking referred for local pain.

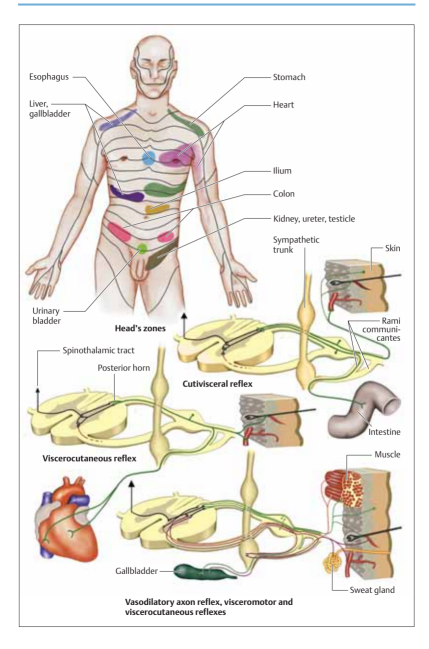
Spinal Autonomic Reflexes

The afferent arm of these reflexes originates in the internal organs and terminates on the sympathetic preganglionic neurons in the intermediolateral and intermediomedial cell columns of the spinal cord at levels T1 through L2 (p. 140). Typical examples are the viscerovisceral reflex (causing meteorism in colic and anuria in myocardial infarction), the viscerocutaneous reflex (a visceral stimulus leads to sweating and hyperemia in the corresponding zone of Head), the cutivisceral reflex (reduction of colic, myogelosis, etc., by warm compresses or massage), the visceromotor reflex (defensive muscle contraction in response to visceral stimulus), and the vasodilatory axon reflex (dermographism). Any abnormality of these reflexes may be an important sign of impaired autonomic function (cardiovascular, gastrointestinal, thermoregulatory, or urogenital), particularly in patients with spinal cord disorders.

Complex Regional Pain Syndrome (CRPS)

The International Association for the Study of Pain (IASP) recommends the term CRPS for a set of painful disorders of apparently related pathophysiology, which are further classified into CRPS type I (reflex sympathetic dystrophy; without peripheral nerve injury) and CRPS type II (causalgia; with peripheral nerve injury). CRPS usually results from a traumatic or other injury to a limb, often in conjunction with prolonged disuse. The pain is persistent and diffuse, and of burning, stabbing, or throbbing quality, often in association with allodynia (pain evoked by a normally nonpainful stimulus) and hyperpathia (abnormally intense pain evoked by a normally painful stimulus). It is generally not in a radicular or peripheral nerve distribution. It may be accompanied by motor disturbances (paresis, disuse of limb), autonomic disturbances (sweat secretion or circulatory disturbances), trophic changes (edema, muscle atrophy, joint swelling, bone destruction), and reactive mental changes (depression, anxiety). The diagnosis of CRPS is based on criteria defined by the IASP and requires the exclusion of other disease processes such as fracture, vasculitis, thrombosis, radicular lesion, rheumatoid arthritis, etc. Its pathogenetic mechanism is unknown

Pain



Circadian Rhythm

The human sleep-wake cycle has a period of approximately 24 hours, as the term *circadian* (Latin *circa* + *dies*) implies. If all external time indicators are removed, the circadian rhythm persists but the times of waking and going to sleep become later each day. The circadian rhythm is thought to be regulated by the suprachiasmatic nucleus of the hypothalamus (p. 142). Retino-hypothalamic connections tie the circadian rhythm to environmental light conditions. There is also a retinal projection to the pineal gland; the melatonin produced there has a rhythm-shifting effect.

Not only sleeping and waking but also many other bodily functions, including cardiovascular and respiratory function, hormone secretion, mitosis rate, intracranial pressure, and attentiveness, follow a circadian pattern (*chronobiology*). Circadian variation in performance is important in the workplace and elsewhere. Some diseases are associated with certain times of the day (*chronopathology*)—certain types of epileptic seizures, asthma, cluster headache, gastroesophageal reflux disease, myocardial infarction, vertricular tachycardia.

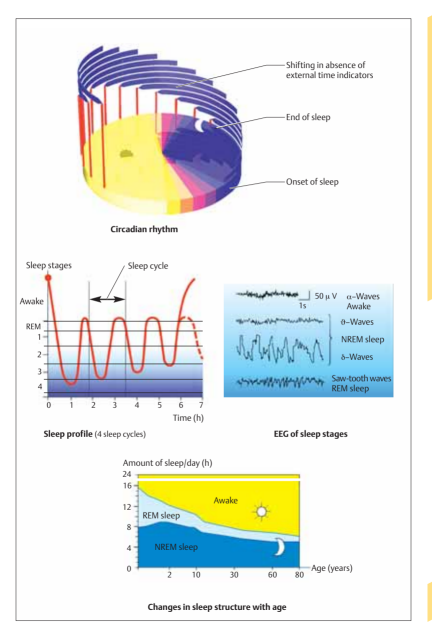
Sleep

Sleep is divided into REM sleep, in which rapid eye movements occur, and non-REM (NREM) sleep. *Polygraphic recordings* (EEG, EOG, EMG) can distinguish these two types of sleep and are used to subdivide NREM sleep into four stages, the last two of which constitute deep sleep (see Table 10, p. 363).

Normal sleep occurs in cycles lasting 90–120 minutes, of which there are thus four or five during a normal night's sleep of ca. 8 hours' duration. Sleep cycles are regulated by activating and deactivating systems (cholinergic REM-on neurons, noradrenergic REM-off neurons) located mainly in the brain stem. The exact physiological significance of sleep is not known. Sleep appears to play a role in regenerative metabolic processes, cognitive functions, and memory.

Sleep Profile

The sleep-wake rhythm changes with age. Neonates sleep 16-18 hours a day at irregular intervals. By age 1 year, the sleep pattern stabilizes to roughly 12 hours of sleep alternating with 12 hours of waking. Adults sleep for 4-10 hours nightly, with the median value ca. 8 hours. As adults age, they tend to take longer and more frequent naps, sleep less deeply, and lie in bed longer in the morning. The sleep architecture changes with age: neonates have 50% REM sleep, but adults only 18%. After age 50, stages 3 and 4 account for only about 5% of sleep. Persons differ in their sleep-wake patterns (somnotypes): there are morning types ("larks") and night types ("night owls"); bedtimes vary by two or more hours among these individuals.



More than 100 sleep disorders have been described to date. Sleep disorders may involve insufficient, interrupted, or absent sleep (*insomnia*), or an excessive need for sleep, including during the day (*hypersonnia*). They may involve respiratory disturbances during sleep (snoring, sleep apnea, asthma), involuntary movement disorders, parasomnias, pharmacologically active substances (medications, illegal drugs, alcohol, coffee, smoking), or systemic disease.

The prerequisite to treatment is an adequate diagnostic assessment, which begins with the determination whether the sleep disorder is primary or secondary (i.e., the result of another disease or condition).

Primary Sleep Disorders (Dyssomnias)

Intrinsic sleep disorders. *Psychogenic insomnia* is characterized by increased mental tension (inability to relax, anxiety, brooding) and excessive concern about sleep itself (constant complaining about an inability to fall asleep or stay asleep, or about waking up too early). Sleep often improves in a new environment (e.g., on vacation).

Pseudoinsomnia is a subjective feeling of disturbed sleep in the absence of objective evidence (i.e., normal polysomnography).

Restless legs syndrome (RLS) is characterized by ascending abnormal sensations in the legs when they are at rest (e.g., when the patient watches television, or before falling asleep) accompanied by an urge to move the legs. It is sometimes present as a genetic disorder with autosomal dominant inheritance. Periodic leg movements during sleep are repeated, abrupt twitching movements of the legs that may persist for minutes to hours. These two movement disorders may appear together or in isolation; both may be either primary or secondary (due to, e.g., uremia, tricyclic antidepressant use, or iron deficiency). Narcolepsy is characterized by daytime somnolence and frequent, sudden, uncontrollable episodes of sleep (imperative sleep), which tend to occur in restful situations (e.g., reading, hearing a lecture, watching TV, long automobile rides). It may be associated with cataplexy (sudden, episodic loss of muscle tone without unconsciousness), sleep paralysis (inability to

move or speak when awaking from sleep), and hypnagogic hallucinations (visual or acoustic hallucinations while falling asleep). Polysomnography reveals a short sleep latency and an early onset of REM sleep. The presence of HLA antigens (DR2, DQw1, DQB1*0602) is nonspecific, as is the absence of hypocretin-1 (orexin A) in the cerebrospinal fluid.

Obstructive sleep apnea is characterized by daytime somnolence with frequent dozing, nocturnal respiratory pauses, and loud snoring. Impaired concentration, decreased performance, and headaches are also common.

Extrinsic sleep disorders. Sleep may be disturbed by external factors such as noise, light, mental stress, and medication use.

Disturbance of the circadian rhythm. Sleep may be disturbed by shift work at night or by intercontinental travel (*jet lag*).

Parasomnias. These disorders include confusion on awakening (sleep drunkenness), sleepwalking (somnambulism), nightmares, sleep myoclonus, bedwetting (enuresis), and nocturnal grinding of the teeth (bruxism).

Secondary Sleep Disorders

Psychogenic sleep disorders. Depression (of various types) can impair sleep, though paradoxically sleep deprivation can ameliorate depression. Depressed persons typically complain of early morning awakening, nocturnal restlessness, and difficulty in starting the day. Sleep disturbances are also common in patients suffering from psychosis, mania, anxiety disorders, alcoholism, and drug abuse.

Neurogenic sleep disorders. Sleep can be impaired by dementia, Parkinson disease, dystonia, respiratory disturbances secondary to neuromuscular disease (muscular dystrophy, amyotrophic lateral sclerosis), epilepsy (nocturnal attacks), and headache syndromes (cluster headaches, migraine). Fatal familial insomnia is a genetic disorder of autosomal dominant inheritance (p. 252).

Sleep disorders due to systemic disease. Sleep can be impaired by pulmonary diseases (asthma, COPD), angina pectoris, nocturia, fibro-myalgia, and chronic fatigue syndrome.

Sleep Disorders



Consciousness is an active process with multiple individual components, including wakefulness, arousal, perception of oneself and the environment, attention, memory, motivation, speech, mood, abstract/logical thinking, and goaldirected action. Psychologists and philosophers have long sought to understand the nature of consciousness.

Clinical assessment of consciousness tests the patients' perception of themselves and their environment, behavior, and responses to external stimuli. Findings are expressed in terms of three categories: level of consciousness (state/clarity of consciousness, quantitative level of consciousness, vigilance, alertness, arousability); content of consciousness (quality of consciousness, awareness); and wakefulness. Changes in any of these categories tend to affect the others as well. Morphologically, the level of consciousness is associated with the reticular activating system (RAS). This network is found along the entire length of the brain stem reticular formation (p. 26), from the medulla to the intralaminar nuclei of the thalamus. The RAS has extensive bilateral projections to the cerebral cortex: the cortex also projects back to the RAS. Neurotransmission in these systems is predominantly with acetylcholine, monoamines (norepinephrine, dopamine, serotonin), GABA (inhibitory), and glutamate (excitatory).

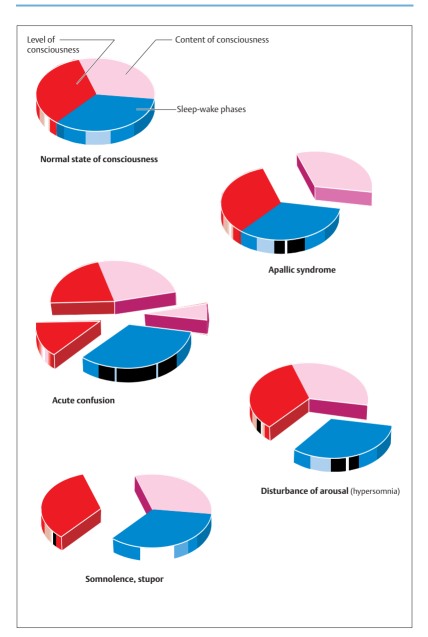
In the normal state of consciousness, the individual is fully conscious, oriented, and awake. All of these categories undergo circadian variation (depending on the time of day, a person may be fully awake or drowsy, more or less concentrated, with organized or disorganized thinking), but normal consciousness with full wakefulness can always be restored by a vigorous stimulus.

Acute Disturbances of Consciousness

Confusion affects the content of consciousness– attention, concentration, thought, memory, spatiotemporal orientation, and perception (lack of recognition). It may also be associated with changes in the level of consciousness (fluctuation between agitation and somnolence) and in wakefulness (impaired sleep–wake cycle with nocturnal agitation and daytime somnolence). *Delirium* is characterized by visual hallucinations, restlessness, suggestibility, and autonomic disturbances (tachycardia, blood pressure fluctuations, hyperhidrosis).

Somnolence is a mild reduction of the *level of* consciousness (drowsiness, reduced spontaneous movement, psychomotor sluggishness, and delayed response to verbal stimuli) while the patient remains arousable: he or she is easily awakened by a stimulus, but falls back asleep once it is removed. The patient responds to noxious stimuli with direct and goal-directed defensive behavior. Orientation and attention are mildly impaired but improve on stimulation. **Stupor** is a significant reduction of the *level* of consciousness. These patients require vigorous and repeated stimulation before they open their eyes and look at the examiner. They answer questions slowly and inadequately, or not at all. They may lie motionless or display restless or stereotyped movements. Confusion reflects concomitant impairment of the content of consciousness.

Disorders of arousal. Wakefulness normally follows a circadian rhythm (p. 112). Sleep apnea syndrome, narcolepsy, and parasomnia are disorders of arousal (dyssomnias, p. 114). *Hypersomnia* is caused by bilateral paramedian thalamic infarcts, tumors in the third ventricular region, and lesions of the midbrain tegmentum (p. 70 ff). The level and content of consciousness may also be affected. In patients with bilateral paramedian thalamic infarction, for example, there may be a sudden onset of confusion, followed by somnolence and coma. After recovery from the acute phase, these patients are apathetic and their memory is impaired ("thalamic dementia").



Coma (from the Greek for "deep sleep") is a state of unconsciousness in which the individual lies motionless, with eves closed, and cannot be aroused even by vigorous stimulation. Coma reflects a loss of the structural or functional integrity of the RAS (p. 116) or the areas to which it projects. Coma may be produced by an extensive brain stem lesion or by extensive bihemispheric cerebral lesions, as well as by metabolic, hypoxic/ischemic, toxic, or endocrine disturbances. In the syndrome of transtentorial herniation (see p. 162), a large unihemispheric lesion can cause coma by compressing the midbrain and the diencephalic RAS. Even without herniation, however, large unihemispheric lesions can transiently impair consciousness.

Coma Staging

The degree of impairment of consciousness is correlated with the extent of the causative lesion. The severity and prognosis of coma are judged from the patient's response to stimuli. There is no universally accepted grading system for coma. Proper documentation involves an exact description of the stimuli given and the responses elicited, rather than isolated items of information such as "somnolent" or "GCS 10." Coma scales (e.g., the Glasgow Coma Scale) are useful for the standardization of data for statistical purposes but do not replace a detailed documentation of the state of consciousness.

Spontaneous movement. Assessment of motor function yields clues to the site of the lesion (p. 44 ff) and the etiology of coma. The examiner should note the pattern of breathing, any utterances, yawning, swallowing, coughing, and movements of the limbs (twitching of the face or hands may indicate epileptic activity; there may be myoclonus or flexion/extension movements).

Stimuli. Lesions of the mid brain or lower diencephalon produce the *decerebration syndrome* (arm/leg extension with adduction and internal rotation of the arms, pronation and flexion of the hands), while extensive bilateral lesions at higher levels produce the *decortication syndrome* (arm/hand flexion, arm supination, leg extension) (p. 47). These pathological flexion and extension movements occur spontaneously or in response to external stimuli (verbal stimulation, tickling around the nose, pressure on the knuckles or other bones) whether the cause of coma is structural or metabolic. Withdrawal of the limb from the stimulus usually means that the pyramidal pathway for the affected limb is intact. Stereotyped flexion or extension movements are usually seen in patients with severe damage to the pyramidal tract.

Brain stem reflexes (p. 26). Structural lesions of the brain stem usually impair the function of the internal and external eye muscles (p. 70 ff), while supratentorial lesions generally do not, unless they secondarily affect the brain stem. Coma in a patient with intact brain stem reflexes is likely to be due to severe bihemispheric dysfunction (if no further objective deficit is found, coma may be psychogenic or factitious; see p. 120). Physicians should be aware that coma due to intoxication or drug overdose (p. 92) may be difficult to distinguish from that due to structural damage by clinical examination alone. Preservation of the vestibulo-ocular reflex (VOR) and of the doll's eyes reflex is compatible with either a bihemispheric lesion or a toxic or metabolic disorder. The VOR induces conjugate eve movement only if its brain stem pathway is intact (from the cervical spinal cord to the oculomotor nucleus). Nonetheless, the VOR may be absent in some cases of toxic coma (due to, e.g., alcohol, barbiturates, phenytoin, pancuronium, or tricyclic antidepressants).

Abnormalities of the respiratory pattern (p. 151) are of limited localizing value. *Cheyne–Stokes respiration* is characterized by regular waxing and waning of the tidal volume, punctuated by apneic pauses. It has a number of causes, including bihemispheric lesions and metabolic disorders. Slow, shallow respiration usually reflects a metabolic or toxic disorder. Rapid, deep respiration (*Kussmaul's respiration*) usually reflects a pontine or mid brain lesion, or metabolic acidosis. Medullary lesions and extensive supratentorial damage produce ataxic, cluster, or gasping respiration.

	Mar and Mar	Absent		0	Absent			
		Flexion/ extension			Absent			ées
	A BANK	Decerebration		0	Sluggish or absent			Diminishing responses and reflexes Stages of coma
N-JEO	A REAL	Decortication	0	000	Sluggish			Diminishin
and the second s	May -	Directed	0	000	Delayed			
MAPLED	K CONTRACTOR	Specifically localized	0	00	Immediate			Normal
Spontaneous movements	Motor response (defensive response) to sensory stimulus		Pupillary diameter	Pupillary light reflex (direct and indirect)		Vestibulo-ocular reflex (doll's -eyes reflex)	Vestibulo-ocular reflex (cold water in either ear; test in left ear shown)	

Coma

Disturbances of Consciousness

119

Comalike Syndromes

Locked-in syndrome (p. 359) is a "de-efferented state" in which the patient is fully conscious but can make no spontaneous movements except lid and vertical eye movements. There may be reflex extension of the arms and legs in response to mild stimuli such as repositioning in bed or tracheal suction. Physicians and nurses must remember that these patients can perceive themselves and their surroundings fully even though they may be unable to communicate. Possible causes include basilar artery occlusion, head trauma, pontine hemorrhage, central pontine myelinolysis, and brain stem encephalitis; a similar clinical picture may be produced by myasthenia gravis, Guillain-Barré syndrome, or periodic paralysis (see pp. 326, 338).

Persistent vegetative state (apallic syndrome) is caused by extensive injury to the cerebral cortex, subcortical white matter, or thalamus. The patients are awake but unconscious (loss of cortical function). Periods in which the eyes are open and move spontaneously, in conjugate fashion, seemingly with fixation, alternate with a state resembling sleep (eyes closed, regular breathing). The patient may blink in response to visual stimuli (rapid hand movements, light), perhaps creating the impression of conscious perception, but does not obey verbal commands. The limbs may be held in a decorticate or decerebrate posture (p. 46). There may be nondirected movements of the arms, legs, head, and jaw, as well as utterances, sucking movements, and lip-licking. The patient may also yawn spontaneously or in response to perioral stimuli. Autonomic disturbances include profuse sweating, tachycardia, urinary and fecal incontinence, and hyperventilation. Optokinetic nystagmus is absent, but the vestibulo-ocular reflex can often be elicited. Spontaneous respiration is preserved. Swallowing is usually possible, but food is kept in the mouth so long than no effective oral nutrition is possible. The persistent vegetative state confers a high mortality. When it lasts for more than a year, improvement is unlikely.

Akinetic mutism. In this syndrome, the patient is awake but the drive to voluntary movement is severely impaired and the patient does not speak (mutism). External stimuli evoke no more than brief ocular fixation without head movement. Possible causes include bifrontal lesions, hydrocephalus, and lesions of the cingulate gyrus or in the third ventricular region. One should keep in mind that other diseases, among them Guillain– Barré syndrome, amyotrophic lateral sclerosis, periodic paralysis, and myasthenia gravis, can present with akinetic mutism or with a similar but less severe syndrome called *abulia* (reduced drive, sluggish voluntary movements, reduced verbal response).

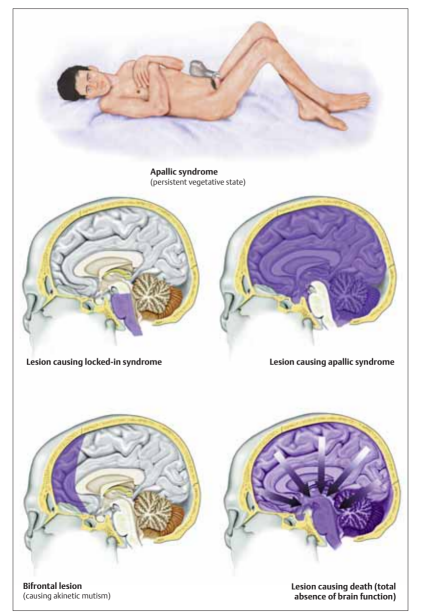
Psychogenic disturbances of consciousness are relatively rare and difficult to diagnose. The lack of arousability can be either an expression of a psychiatric disease (conversion or acute stress reaction, severe depression, catatonic stupor) or a deliberate fabrication. Clues are sometimes found in the case history or on neurological examination (e. g., presence of aversive reflexes, active eye closing, preserved optokinetic and vestibulo-ocular nystagmus, catalepsy, stereotyped posture).

Death

Death is medically and legally defined as the total and irreversible cessation of all brain function (hence the synonymous term, "brain death"). Spontaneous respiration (a function of the brain stem) is absent, though the heart may continue beating and other organs may still function if supportive measures are maintained (ventilation, pressor medications). All organ systems cease to function when these are discontinued.

The clinical determination of death is based on the following criteria: coma; lack of spontaneous respiration (apnea test); lack of response to noxious stimuli (with the possible exception of spinal reflexes); absence of brain stem reflexes (pupillary, corneal, cough, gag, and oculovestibular reflexes). The diagnosis of death requires the exclusion of possibly similarappearing states such as toxic, metabolic, and endocrine disorders, pharmacological relaxation and sedation, and hypothermia. Major structural damage of the brain is present in all cases (though not necessarily demonstrable on all imaging studies). Ancillary diagnostic testing (EEG, Doppler sonography, evoked potentials, perfusion scintigraphy, cerebral angiography, MRI) may support the diagnosis but is generally not legally required (Table 11, p. 364).

Disturbances of Consciousness



Personality is the set of physical and psychological traits that distinguish one individual from another. It evolves over time under the influence of changes in brain function, as well as other internal and external factors. These include neurobiological factors (heredity, structure and function of the nervous system), physiological factors (endocrine, metabolic), socialization (formation of language, thought, emotion, and action according to societal norms and value systems), individualization (consciousness of one's own individuality), sexuality, temperament, intelligence, life experiences, education, economic status, and individual will. Changes in personality cause gradual or abrupt changes in behavior. Some neurological diseases produce behavioral changes; the clinical picture depends mainly on the location of the disturbance.

Frontal Lobe Lesions

The frontal lobe includes the motor cortex (areas 4, 6, 8, 44), the prefrontal cortex (areas 9–12 and 45–47), and the cingulate gyrus (p. 144). It is responsible for the planning, monitoring, and performance of motor, cognitive, and emotional functions (*executive functions*). Frontal lobe syndromes may be due to either cortical or subcortical damage and thus cannot be reliably localized without neuroimaging. The typical syndromes listed here are useful for classification but do not imply a specific diagnosis or exact localization of the underlying lesion.

Lateralized syndromes. Left frontal lobe lesions, depending on their location and extent, can produce right hemiparesis or hemiplegia, transcortical motor aphasia and diminished verbal output (p. 126), buccofacial apraxia (p. 128), and/or depression or anxiety. *Right frontal lobe lesions* can produce left hemiparesis or hemiplegia, left hemineglect (p. 132), mania, and/or increased psychomotor activity.

Nonlateralized syndromes. Fronto-orbital lesions produce increased drive, memory impairment with confabulation, and disorientation. Disinhibition and impaired insight into one's own behavior may produce abnormal facetiousness (German Witzelsucht), abnormal social behavior (loss of distance, sexual impulsiveness), indifference, or carelessness. Lesions of the *cingulate gyrus* and *premotor cortex* produce syndromes ranging from abulia (loss of drive) to akinetic mutism (p. 120) and generally characterized by apathy, loss of interest, inertia, loss of initiative, decreased sexual activity, loss of emotion, and loss of planning ability. Urinary and fecal incontinence occur because of the loss of (cortical) perception of the urge to urinate and defecate. Altered voiding frequency or sudden voiding is the result.

These patients are usually impaired in their capacity for *divided attention* (the processing of new information and adaptation to altered requirements, i.e., flexibility) and for *directed attention* (selective attention to a particular thing or task). Their attention span is short, they are easily distracted, they have difficulty in the execution of motor sequences, and they tend to perseverate (to persist in a particular activity or thought). Increased distractibility and prolonged reaction times impair performance in the workplace and in everyday activities such as driving.

Lesions of pathways. Lesions in pathways connecting the frontal lobe to other cortical and subcortical areas (p. 24) can produce frontal lobe-type syndromes, as can other diseases including multisystem atrophy, Parkinson disease, Alzheimer disease, normal-pressure hydrocephalus, and progressive supranuclear palsy. Lesions of the corpus callosum. See p. 24.

Behavioral Changes



Abulia



Anxiety, misperceptions



Concentration and attention deficits



Pathological crying and laughing

Defensiveness, irritability, psychomotor agitation

Language is a means of transmitting and processing information, organizing sensory perceptions, and expressing thoughts, feelings, and intentions. The content of language encompasses the past, present, and future. The development of language does not necessarily require speech and audition: deaf-mutes learn to communicate with sign language. Language is most easily acquired in childhood. Linguistic messages are transmitted and received through speaking and hearing, writing and reading, or (in the case of sign language) the production and interpretation of gestures. The cerebral language areas are located in the left hemisphere in over 90% of right-handers and in 60% of left-handers; the remaining individuals have bihemispheric or (in 1-2%) exclusively right-hemispheric dominance for language. The left (dominant) hemisphere is responsible for the cognitive processing of language, while the right (nondominant) hemisphere produces and recognizes the emotional components of language (prosody = emphasis, rhythm, melody). Language is subserved by subcortical nuclei as well (left thalamus, left caudate nucleus, associated fiber pathways). Language function depends on the well-coordinated activity of an extensive neural network in the left hemisphere. It is simplistic to suppose that language is understood and produced by means of a unidirectional flow of information through a chain of independently operating brain areas linked together in series. Rather, it has been shown that any particular linguistic function (such as reading, hearing, or speaking) relies on the simultaneous activation of multiple, disparate cortical areas. Yet the simplified model of language outlined below (proposed by Wernicke and further elaborated by Geschwind) usually suffices for the purposes of clinical diagnosis.

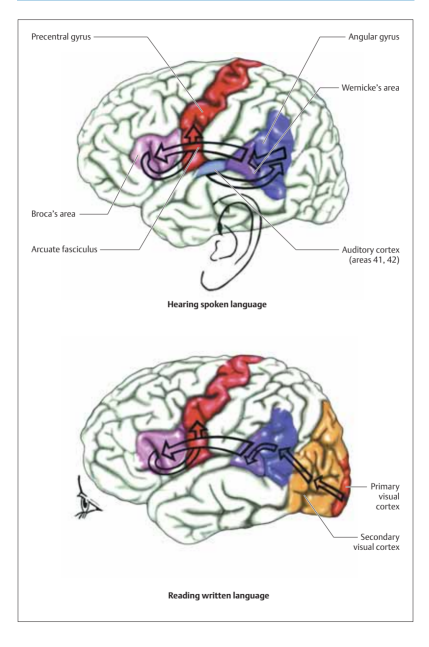
Hearing and speaking. Acoustic signals are transduced in the inner ear into neural impulses in the cochlear nerve, which ascend through the auditory pathway and its relay stations to the primary and secondary auditory cortex (p. 100). From here, the information is sent to *Wernicke's area* (the "posterior language area"), consisting of Wernicke's area proper, in the superior temporal gyrus (Brodmann area 22), as well as the angular and supramarginal gyri (areas 39, 40). The *angular gyrus* processes auditory, visual,

and tactile information, while Wernicke's area proper is the center for the understanding of language. It is from here that the arcuate fasciculus arises, the fiber tract that conveys linguistic information onward to *Broca's area* (areas 44 and 45; the "anterior language area"). Grammatical structures and articulation programs are represented in Broca's area, which sends its output to the motor cortex (speech, p. 130). Spoken language is regulated by an auditory feedback circuit in which the utterer hears his or her own words and the cortical language areas modulate the speech output accordingly.

Reading and writing. The visual pathway conveys visual information to the primary and secondary visual cortex (p. 80), which, in turn, project to the angular gyrus and Wernicke's area, in which visually acquired words are understood, perhaps after a prior "conversion" to phonetic form. Wernicke's area then projects via the arcuate fasciculus to Broca's area, as discussed above; Broca's area sends its output to the motor cortex (for speech or, perhaps, to the motor hand area for writing). This pathway enables the recognition and comprehension of written language, as well as reading out loud.

Examination. The clinical examination of language includes spontaneous speech, naming of objects, speech comprehension, speech repetition, reading, and writing. The detailed assessment of aphasia requires the use of test instruments such as the Aachen aphasia test, perhaps in collaboration with neuropsychologists and speech therapists. Disturbances of speech may be classified as *fluent* or *nonfluent*. Examples of the former are paragrammatism (faulty sentence structure), meaningless phrases, circumlocution, semantic paraphasia (contextual substitution, e.g., "leg" for "arm"), phonemic paraphasia (substitution of one letter for another, e.g., "tan" for "can"), neologisms (nonexistent words), and fluent gibberish (jargon). Examples of the latter are agrammatism (word chains without grammatical structure), echolalia (repetition of heard words), and automatism (repeating the same word many times). Prosody and dysarthria (if present; p. 130) are evaluated during spontaneous speech. Anomia is the inability to name objects. Patients with aphemia can read, write, and understand spoken language but cannot speak.

Language



Aphasia

Aphasia is an acquired disturbance of language. Lesions at various sites produce different types of aphasia; focal lesions do not cause total loss of all language functions simultaneously. The side of cerebral dominance for language can be determined by the Wada test (intracarotid amobarbital procedure, IAP), in which amobarbital is injected first into one internal carotid artery and then into the other, under angiographic control, to selectively anesthetize each hemisphere (this is done, for example, before cortical resections for epilepsy). Crossed aphasia, i.e., aphasia due to a right hemispheric lesion in a right-handed patient, is rare. Aphasia usually improves markedly within a few weeks of onset and may continue to improve gradually over the first year, even if the symptoms temporarily appear to have stabilized. Improvement beyond one year is rare and usually minor.

Aphasia in bilingual and multilingual persons (usually) affects all of the languages spoken. The severity of involvement of each language depends on the age at which it was acquired, premorbid language ability, and whether the languages were learned simultaneously or sequentiallly. Aphasia is most commonly due to stroke or head trauma and may be accompanied by apraxia.

Global aphasia involves all aspects of language and severely impairs spoken communication. The patient cannot speak spontaneously or can only do so with great effort, producing no more than fragments of words. Speech comprehension is usually absent; at best, patients may recognize a few words, including their own name. Perseveration (persistent repetition of a single word/subject) and neologisms are prominent, and the ability to repeat heard words is markedly impaired. Patients have great difficulty naming objects, reading, writing, and copying letters or words. Their ability to name objects, read, and write, except for the ability to copy letters of the alphabet or isolated words, is greatly impaired. Language automatism (repetition of gibberish) is a characteristic feature. Site of lesion: Entire distribution of the middle cerebral artery, including both Broca's and Wernicke's areas.

Broca's aphasia (also called anterior, motor, or expressive aphasia) is characterized by the absence or severe impairment of spontaneous speech, while comprehension is only mildly impaired. The patient can speak only with great effort, producing only faltering, nonfluent, garbled words. Phonemic paraphasic errors are made, and sentences are of simple construction, often with isolated words that are not grammatically linked (agrammatism, "telegraphic" speech). Naming, repetition, reading out loud, and writing are also impaired. Site of lesion: Broca area; may be due to infarction in the distribution of the prerolandic artery (artery of the precentral sulcus). Wernicke's aphasia (also called posterior, sensory, or receptive aphasia) is characterized by severe impairment of comprehension. Spontaneous speech remains fluent and normally paced, but paragrammatism, paraphasia, and neologisms make the patient's speech partially or totally incomprehensible (word salad, jargon aphasia). Naming, repetition of heard words, reading, and writing are also markedly impaired. Site of lesion: Wernicke's area (area 22). May be due to infarction in the distribution of the posterior temporal artery.

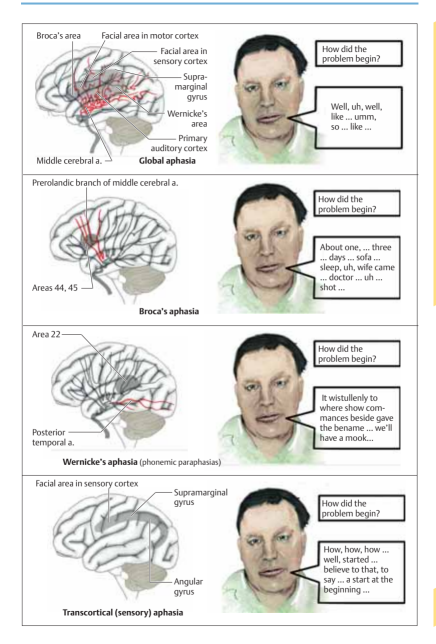
Transcortical aphasia. Heard words can be repeated, but other linguistic functions are impaired: spontaneous speech in transcortical *motor* aphasia (syndrome similar to Broca's aphasia), language comprehension in transcortical *sensory* aphasia (syndrome similar to Wernicke's aphasia). *Site of lesion:* Motor type, left frontal lobe bordering on Broca's area; sensory type, left temporo-occipital junction dorsal to Wernicke's area. Watershed infarction is the most common cause (p. 172).

Amnestic (anomic) aphasia. This type of aphasia is characterized by impaired naming and wordfinding. Spontaneous speech is fluent but permeated with word-finding difficulty and paraphrasing. The ability to repeat, comprehend, and write words is essentially normal. *Site of lesion:* Temporoparietal cortex or subcortical white matter.

Conduction aphasia. Repetition is severely impaired; fluent, spontaneous speech is interrupted by pauses to search for words and by phonemic paraphasia. Language comprehension is only mildly impaired. *Site of lesion:* Arcuate fasciculus or insular region.

Subcortical aphasia. Types of aphasia similar to those described may be produced by subcortical lesions at various sites (thalamus, internal capsule, anterior striatum).

Aphasia



Agraphia. Agraphia is the acquired inability to write. Agraphia may be isolated (due to a lesion located in area 6, the superior parietal lobule, or elsewhere) or accompanied by other disturbances: aphasic agraphia is fluent or nonfluent, depending on the accompanying aphasia; apraxic agraphia is due to a lesion of the dominant parietal lobe; spatial agraphia, in which the patient has difficulty writing on a line and only writes on the right side of the paper, is due to a lesion of the nondominant parietal lobe; alexia with agraphia may be seen in the absence of aphasia. Micrographia (abnormally small handwriting) is found in Parkinson disease (p. 206) and is not pathogenetically related to agraphia. Various forms of agraphia are common in Alzheimer disease. Examination: The patient is asked to write sentences, long words, or series of numbers to dictation, to spell words, and to copy written words.

Alexia. Alexia is the acquired inability to read. In isolated alexia (alexia without agraphia), the patient cannot recognize entire words or read them quickly, but can decipher them letter by letter, and can understand verbally spelled words. The ability to write is unaffected. The responsible lesion is typically in the left temporooccipital region with involvement of the visual pathway and of callosal fibers. Anterior alexia (difficulty and errors in reading aloud; impaired ability to write, spell, and copy words) is usually associated with Broca's aphasia. Central alexia (combination of alexia and agraphia) is usually accompanied by right-left disorientation, finger agnosia, agraphia, and acalculia (Gerstmann syndrome; lesions of the angular and supramarginal gyri), or by Wernicke's aphasia. Other features include the inability to understand written language or to spell, write, or copy words. Examination: The patient is asked to read aloud and to read individual words, letters, and numbers; the understanding of spelled words and instructions is tested.

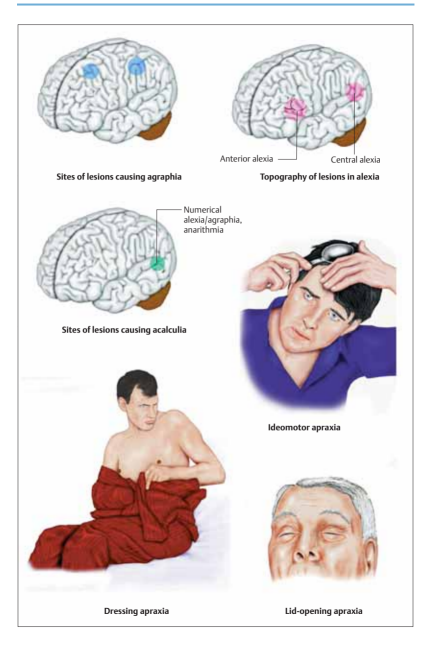
Acalculia. Acalculia is an acquired inability to use numbers or perform simple arithmetical calculations. Patients have difficulty counting change, using a thermometer, or filling out a check. Lesions of various types may cause acalculia. *Examination:* The patient is asked to perform simple arithmetical calculations and to read numbers. **Apraxia.** There are several kinds of apraxia; in general, the term refers to the inability to carry out learned motor tasks or purposeful movements. Apraxia is often accompanied by aphasia.

Ideomotor apraxia involves the faulty execution (parapraxia) of acquired voluntary and complex movement sequences; it can be demonstrated most clearly by asking the patient to perform pantomimic gestures. It can involve the face (buccofacial apraxia) or the limbs (limb apraxia). It is due to a lesion in the association fiber pathways connecting the language, visual, and motor areas to each other and to the two hemispheres (disconnection syndrome). Examination (pantomimic gestures on command): face (open eyes, stick out tongue, lick lips, blow out a match, pucker, suck on a straw); arms (turn a screw, cut paper, throw ball, comb hair, brush teeth, snap fingers); legs (kick ball, stamp out cigarette, climb stairs). The patient may perform the movement in incorrect sequence, or may carry out a movement of the wrong type (e.g., puffing instead of sucking).

Ideational apraxia is impairment of the ability to carry out complex, learned, goal-directed activities in proper logical sequence. A temporal or parietal lesion may be responsible. *Examination:* The patient is asked to carry out pantomimic gestures such as opening a letter, making a sandwich, or preparing a cup of tea.

Apraxia-like syndromes. The following disturbances are termed "apraxia" even though actual parapraxia is absent: *Lid-opening apraxia* (p. 64) is difficulty opening the eyes on command. *Gait apraxia* is characterized by difficulty initiating gait and by short steps (p. 160). *Dressing apraxia* is often seen in patients with nondominant parietal lobe lesions. They cannot dress themselves and do not know how to position a shirt, shoes, trousers, or other items of clothing to put them on correctly. An underlying impairment of spatial orientation is responsible.





Speech

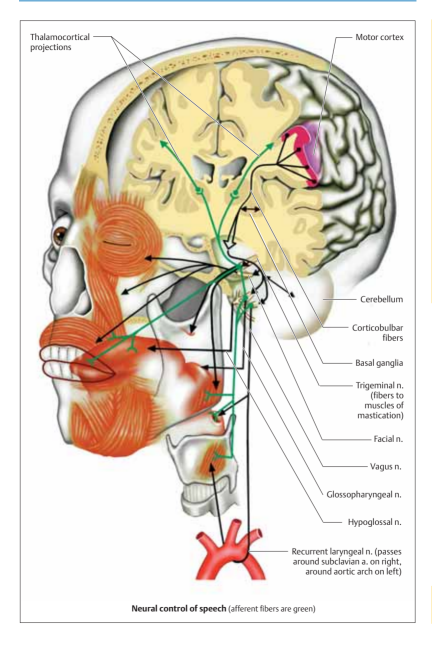
The neural basis of speech. Speech-related movement programs generated in the premotor cortex (area 6) are modulated by information from the cerebellum and basal ganglia and are relayed to the motor cortex (inferior portion of the precentral gyrus, area 4) for implementation. The motor cortex projects by way of the corticopontine and corticobulbar tracts to the motor cranial nerve nuclei in the brain stem. CN V (mandibular nerve) controls the muscles that open and close the jaw (masseter, temporalis, medial and lateral pterygoid muscles). CN VII controls facial expression and labial articulation; CN X and, to a lesser extent, IX control motility of the soft palate, pharynx, and larynx; CN XII controls tongue movement. Speech-related impulses to the respiratory muscles travel (among other pathways) from the motor cortex to the spinal anterior horn cells. Connections to the basal ganglia and cerebellum are important for the coordination of speech. Sensory impulses from the skin, mucous membranes, and muscles return to the brain through CN V (maxillary and mandibular nerves), IX, and X. These impulses are processed by a neural network (reticular formation, thalamus, precentral cortex) mediating feedback control of speech. The central innervation of the speech pathway is predominantly bilateral; thus, dysarthria due to unilateral lesions is usually transient.

Voice production (phonation). Voice production by the larynx (phonation) through the vibrating vocal folds (cords) yields sound at a fundamental frequency with a varying admixture of higher-frequency components, which lend the voice its timbre (musical quality); timbre depends on the resonant cavities above the vocal folds (pharynx, oral cavity, nasal cavity). The volume of the voice is regulated by stretching and relaxation of the vocal folds and by adjustment of air pressure in the larynx. The air flow necessary for phonation is produced in the respiratory tract (diaphragm, lungs, chest, and trachea). The individual structural characteristics of the larynx, particularly the length of the vocal folds, determine the pitch of a person's voice. A whisper is produced when the vocal folds are closely apposed and do not vibrate.

Creation of the sounds of speech (articulation). The sounds of speech are created by changing the configuration of the physiological resonance spaces and articulation zones. The resonance spaces can be altered by movement of the velum (which separates the oral and pharyngeal cavities) and tongue (which divides the oral cavity). Each vowel (a, e, i, o, u) is associated with a specific partitioning of the oral cavity by the tongue. The palate, teeth, and lips are the articulation zones with which consonants are produced (g, s, b, etc.).

Dysarthria, Dysphonia

Dysarthria (impaired articulation) and dysphonia (impaired phonation and resonance) result from a disturbance of the neural control mechanism for speech (sensory portion, motor portion, or both). Diagnostic assessment requires both analysis of the patient's vocal output (breathing, phonation, resonance, articulation; speed, coordination, and prosody of speech) and the determination of any associated neurological findings (e.g. dysphagia, hyperkinesia, cranial nerve deficits). For responsible lesions and syndromes, see Table 12 (p. 365).



Agnosia is defined as a disturbance of recognition in which perception, attention, and general intelligence are (largely) unimpaired.

Disturbances of Body Image Perception

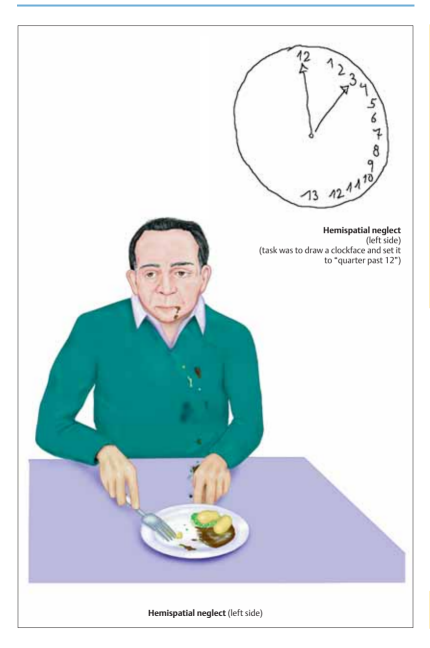
Autotopagnosia (body-image agnosia) is the inability to correctly orient or perceive different body parts; patients cannot obey commands to point to parts of their own or the examiner's body (e.g., foot, hand, nose). The responsible lesion is usually, though not always, in the temporoparietal region (angular and supramarginal gyri). An aphasic patient may appear to have autotopagnosia because he cannot understand verbal instructions, but aphasia may also coexist with true autotopagnosia. Finger agnosia is the inability to identify, name, or point to fingers. These patients cannot mimic the examiner's finger movements or copy finger movements of their own contralateral hidden hand with the affected hand. Right-left disorientation is the inability to distinguish the right and left sides of one's own or another's body; these patients cannot obey a command to raise their left hand or touch it to their right ear. This type of disorientation can cause dressing apraxia (p. 128) and similar problems.

Anosognosia is the unawareness or denial of a neurological deficit, such as hemiplegia. Patients may claim that they only want to give the paralyzed side a rest, or attempt to demonstrate that their condition has improved without realizing that they are moving the limb on the unaffected side. Most such patients have extensive lesions of the nondominant hemisphere. Anosognosia may also accompany visual field defects due to unilateral or bilateral lesions of the visual cortex (homonymous hemianopsia, cortical blindness). The most striking example of this is Anton syndrome, in which cortically blind patients act as if they could see, and will even "describe" details of their surroundings (incorrectly) without hesitation.

Disturbances of Spatial Orientation

A number of different types of agnosia impair the awareness of one's position relative to the surroundings, i.e., *spatial orientation*. Parietooccipital lesions are commonly responsible. Constructional apraxia is characterized by the inability to represent spatial relationships in drawings, or with building blocks. Affected patients cannot copy a picture of a bicycle or clock. Everyday activities are impaired by the inability to draw diagrams, read (analog) clocks, assemble pieces of equipment or tools, or write words in the correct order (spatial agraphia). Hemineglect is the inability to consciously perceive, react to, or classify stimuli on one side in the absence of a sensorimotor deficit or exceeding what one would expect from the severity of the sensorimotor deficit present. Hemineglect may involve unawareness of one side of the body (one-sided tooth brushing, shaving, etc.) or of one side of an object (food may be eaten from only one side of the plate, eyeglasses may be looked for on only one side of the room). When addressed, the patient always turns to the healthy side. Neurological examination reveals that double simultaneous stimulation (touch, finger movement) of homologous body parts (same site, e.g., face or arm) is not felt on the affected side (extinction phenomenon). In addition, perception of stimuli on the affected side is quantitatively lower than on the healthy side, there is limb akinesia despite normal strength on the side of the lesion, and spatial orientation is impaired (e.g., the patient copies only half of a clock-face).

Disturbances of Orientation



Memory

Memory involves the acquisition, storage, recall, and reproduction of information. Memory depends on intact functioning of the limbic system (p. 144) and areas of the brain that are connected to it.

Declarative or explicit memory (i.e., memory for facts and events) can be consciously accessed and depends on intact functioning of the mediobasal portion of the temporal lobe. The duration of information storage may be relatively short (short-term, immediate, and working memory) or long (long-term memory). Verbal (telephone number) or visuospatial information (how to find a street) can be directly recalled from shortterm memory. The entorhinal cortex plays a key role in these memory functions: all information from cortical regions (frontal, temporal, parietal) travels first to the entorhinal cortex and then, by way of the parahippocampal and perirhinal cortex, to the hippocampus. There is also a reciprocal projection from the hippocampus back to the entorhinal cortex. Long-term memory stores events of personal history that occurred at particular times (episodic memory for a conversation, one's wedding day, last year's holiday; orbitofrontal cortex) as well as conceptual, non-time-related knowledge (semantic memory for the capital of Spain, the number of centimeters in a meter, the meaning of the word "stethoscope"; subserved by different cortical regions).

Nondeclarative (procedural, implicit) memory, on the other hand, cannot be consciously accessed. Learned motor programs (riding a bicycle, swimming, playing the piano), problem-solving (rules), recognition of information acquired earlier (priming), and conditioned learning (avoiding a hot burner on the stove, sitting still in school) belong to this category. Nondeclarative memory is mediated by the basal ganglia (motor function), neocortex (priming), cerebellum (conditioning), striatum (agility), amygdala (emotional responses), and reflex pathways.

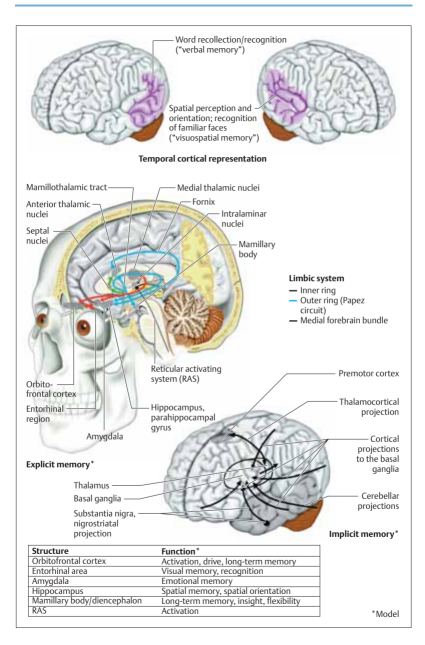
Examination. Only disturbances of declarative memory (*amnesia*) can be studied by clinical examination. *Short-term memory*: the acquisition of new information is tested by having the patient repeat a series of numbers or groups of words and asking for this information again

5–10 minutes later. The patient's *orientation* (name, place of residence/address, time/date) and *long-term memory* (place of birth, education, place of employment, family, general knowledge) are also tested by directed questioning.

Memory Disorders (Amnesia)

Forgetfulness. Verbal memory does not decline until approximately age 60, and even then only gradually, if at all. Aging is, however, often accompanied by an evident decline in information processing ability and attention span (*benign senescent forgetfulness*). These changes occur normally, yet to a degree that varies highly among individuals, and they are often barely measurable. They are far less severe than fullblown dementia, but they may be difficult to distinguish from incipient dementia.

Amnesia. Anterograde amnesia is the inability to acquire (declarative) information, for later recall, from a particular moment onward; *retrograde amnesia* is the inability to remember (declarative) information acquired before a particular moment (p. 269). Amnestic patients commonly *confabulate* (i.e., fill in gaps in memory with fabricated, often implausible information); they may be disoriented and lack awareness of their own memory disorder. For individual symptoms and their causes, see Table 13 (p. 365).



Dementia

Dementia is a newly occurring, persistent, and progressive loss of cognitive function. Both short-term and long-term memory are impaired, in conjunction with at least one of the following disorders: aphasia, apraxia, agnosia, or impairment of abstract thinking, decisionmaking ability. visuospatial performance, planned action, or personality. Professional, social and interpersonal relationships deteriorate, and the sufferer finds it increasingly difficult to cope with everyday life without help. The diagnosis of dementia requires the exclusion of disturbances of consciousness (e.g., delirium) and of psychiatric disease (e.g., depression, schizophrenia). The differential diagnosis also includes benign senescent forgetfulness ("normal aging," in which daily functioning is unimpaired) and amnestic disorders. Approximately 90% of all cases of dementia are caused by Alzheimer disease (p. 297) or cerebrovascular disorders; diverse etiologies account for the rest. The physician confronted with a case of incipient dementia must distinguish primary dementia from that secondary to another disease (Table 14, p. 366). The objective is early determination of the etiology of dementia, especially when these are treatable or reversible.

Examination. The patient or another informant should be asked for an account of the duration, type, and extent of problems that arise in every-

day life. The clinical examination is used to ascertain the type and severity of cognitive deficits and any potential underlying disease. Standardized examining instruments are useful for precise documentation and differentiation of the cognitive deficits. Rapid tests for dementia. such as the Mini-Mental Status Examination. mini-syndrome test, and clock/numbers test, are useful for screening. Function-specific neurophysiological tests permit diagnostic assessment of individual aspects of cognition includorientation. attention. concentration, ing memory, speech, and visual constructive performance. Laboratory tests (ESR1, differential blood count, electrolytes, liver function tests, BUN², creatinine, glucose, vitamin B₁₂, folic acid, TPHA3, TSH4, and HIV5), EEG, and diagnostic imaging techniques (CT⁶, MRI⁷, SPECT⁸, and PET⁹) provide further useful information for classification and determination of the cause of dementia. None of these diagnostic techniques alone can pinpoint the etiology of dementia; definitive diagnosis practically always requires multiple tests and examinations. Diagnostic imaging is of particular importance in patients with the subacute onset of cognitive impairment or amnesia (<1 month), fluctuation or acute worsening of symptoms, papilledema, visual field defects, headaches, a recent head injury, known malignancies, epilepsy, a history of stroke, urinary incontinence, or an abnormal gait.

¹ESR Erythrocyte sedimentation rate

²BUN Blood urea nitrogen

³TPHA Treponema pallidum hemagglutination test

⁴TSH Thyroid-stimulating hormone

⁵HIV Human immunodeficiency virus

⁶CT Computerized tomography

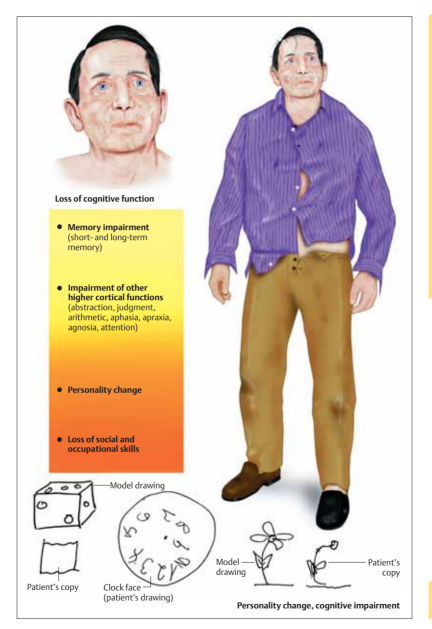
⁷MRI Magnetic resonance imaging

⁸SPECT Single photon emission computerized tomography

⁹PET Positron emission tomography

Additional diagnostic tests to be performed as needed: Coagulation profile, serum protein electrophoresis, serum ammonia, parathyroid hormone, cortisol, rheumatoid factor, antinuclear antibodies, blood alcohol, serum/urine drug levels, copper/ceruloplasmin, lactate/pyruvate, hexosaminidase, CSF tests, molecular genetic analysis.

Dementia



137

Patients with symptoms and signs that are unusual, difficult to classify, or resistant to treatment are often referred to a neurologist for a determination whether the patient's problem is "organic" or "psychogenic." Many such patients make a diagnostic and therapeutic odyssey from one medical or paramedical office to another, and have a long list of positive findings to show for it. In other patients, symptoms and signs may arise acutely or subacutely, perhaps in repeated episodes, creating the impression of a serious illness. The physician's primary objectives must be (1) to identify the possible physical or psychosocial causes of the problem, and at the same time (2) to avoid unnecessary or dangerous diagnostic tests. A correct diagnosis requires time, solid knowledge of the relevant anatomy and physiology, and the ability to recognize the psychosocial dynamics that may have given rise to the patient's complaints.

If detailed neurological examination reveals no abnormality and the symptoms cannot be attributed to any neurological disease, the physician should consider potential psychosocial causes. These may be unconscious (e.g., an inner conflict of which the patient is unaware) or conscious (e.g., a deliberate attempt to acquire the financial benefits and increased attention associated with illness). The underlying cause may be an unresolved social conflict (familial, professional, financial) or some other mental disorder (depression, anxiety, obsessive-compulsive disorder, personality disorder). Organic dysfunction and objective signs of illness are disproportionately mild in relation to the patient's complaints, unrelated to them, or entirely absent.

Conversion disorders (previously termed "conversion hysteria") often present with a single (pseudoneurological) symptom, such as psychogenic amnesia, stupor, mutism, seizures, paralysis, blindness, or sensory loss. It has been theorized that such symptoms serve to resolve unconscious inner conflicts. The diagnosis may be particularly difficult to make in patients who simultaneously suffer from organic neurological or psychiatric disease (e.g., hyperventilation in epilepsy, headaches in depression, paralysis in multiple sclerosis).

Somatoform disorders, according to current psychiatric terminology, are mental disorders

characterized by "repeated presentation of physical symptoms, together with persistent requests for medical investigations, in spite of repeated negative findings and reassurances by doctors that the symptoms have no physical basis" (ICD-10, WHO, 1992). In somatization disorder, the patient asks for treatment of multiple, recurrent, and frequently changing symptoms, which often affect multiple organ systems (e.g. headaches + bladder dysfunction + leg pains + breathing disorder). In hypochondriacal disorder (previously termed "hypochondriasis"), the patient is less concerned about the symptoms themselves, and more preoccupied with the supposed presence of a serious disease. The fears persist despite repeated, thorough examination, normal test results, and medical reassurance. Any mild abnormalities that may happen to be found, e.g. of heartbeat, respiration, or intestinal function, or skin changes, only amplify the patient's anxiety. Persistent somatoform pain disorder involves complaints of "persistent, severe, and distressing pain, which cannot be explained fully by a physiological process or a physical disorder" (ICD-10), though an organic cause of pain is often present as well. The physical impairment that the patient attributes to pain may actually be due to a lack of fulfillment in familial, professional, or social relationships. For these patients, dealing with the pain may become the major "purpose in life." Malingering is not a mental disorder, but rather

the deliberate, premeditated feigning of illness to achieve a goal (e.g., feigning of headaches to obtain opiates).

Simulated or intentionally induced (factitious) symptoms may serve no clear purpose (neither the resolution of an unconscious conflict, nor any obvious kind of gain); they may be present in the simulator (*Münchhausen syndrome*) or in a child or other person under their care (*Münchhausen-by-proxy*). The peregrinating patient (*hospital hopper*) demands diagnostic tests from one physician after another, but negative test results can never put the patient's fears to rest. Patients with *Ganser syndrome* give approximate or fatuous answers to simple questions, possibly creating the impression of dementia.

Behavioral Manifestations of Neurological Disease



Organization

The autonomic nervous system (ANS) is so called because its functions are normally not subject to direct voluntary control. It regulates hormonal and immunological processes as well as the functioning of major organ systems (cardiovascular, respiratory, gastrointestinal, urinary, and reproductive systems).

ANS, Central Portion

Central components of the ANS are found in the cerebral cortex (insular, entorhinal, orbitofrontal, and frontotemporal areas), the hypothalamus, the limbic system, the mid brain (periaqueductal gray substance), the medulla (nucleus of the solitary tract, ventrolateral and ventromedial areas of the medulla), and the spinal cord (various tracts and nuclei, discussed in further detail below).

Afferent connections. Afferent impulses enter the ANS from spinal tracts (anterolateral fasciculus = spinothalamic + spinocerebellar + spinoreticular tracts), brain stem tracts (arising in the reticular formation), and corticothalamic tracts, and from the *circumventricular* organs. The latter are small clusters of specialized neurons, lying on the surface of the ventricular system, that sense changes in the chemical composition of the blood and the cerebrospinal fluid (i.e., on both sides of the blood-CSF barrier). These organs include the organum vasculosum of the lamina terminalis (in the roof of the third ventricle behind the optic chiasm \Rightarrow cytokines/ fever), the subfornical organ (under the fornices between the foramina of Monro ⇒ angiotensin II/blood pressure and fluid balance), and the area postrema (rostral to the obex on each side of the fourth ventricle \Rightarrow cholecystokinin/ gastrointestinal function, food intake).

Efferent connections. Projections from the hypothalamus and brain stem, particularly from the brain stem reticular formation, travel to the lateral horn of the thoracolumbar spinal cord, where they form synapses onto the sympathetic neurons of the spinal cord. The latter, in turn, project preganglionic fibers to the sympathetic ganglia (p. 147). The parasympathetic neurons receive input from higher centers in similar fashion and project in turn to parasympathetic ganglia that are generally located near the end organs they serve. The hypothalamus regulates hormonal function through its regulator hormones as well as efferent neural impulses.

Neurotransmitters. The main excitatory neurotransmitter is *glutamate*, and the main inhibitory neurotransmitter is γ -aminobutyric acid (GABA). Modulating neurotransmitters include acetylcholine, amines, neuropeptides, purines, and nitric oxide (NO).

ANS, Peripheral Portion (p. 146)

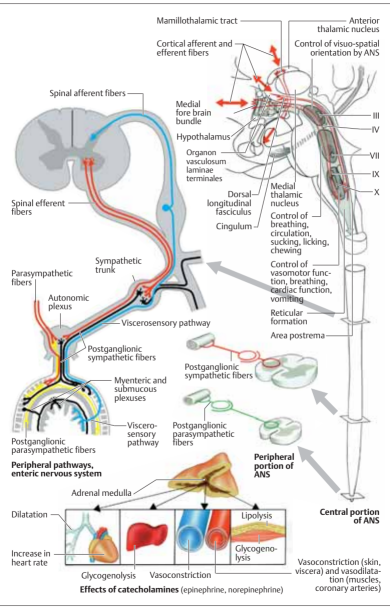
The sympathetic and parasympathetic components are both structurally and functionally segregated.

Spinal nuclei. *Sympathetic* spinal neurons lie in the lateral horn (intermediolateral and intermediomedial cell columns) of the thoracolumbar spinal cord (T1–L2) and are collectively termed the *thoracolumbar system*. *Parasympathetic* spinal neurons lie in the brain stem (with projections along CN III, VII, IX, X) and the sacral spinal cord (S2–S4), and are collectively termed the *craniosacral system*. The intestine has its own autonomic ganglia, which are located in the myenteric and submucous plexueses (p. 154).

Afferent connections. Afferent impulses to the ANS enter the spinal cord via the dorsal roots, and the brain stem via CN III, VII, IX and X.

Efferent connections. The projecting fibers of the spinal autonomic neurons (preganglionic fibers) exit the spinal cord in the ventral roots and travel to the paravertebral and prevertebral ganglia, where they synapse onto the next neuron of the pathway. The sympathetic preganglionic fibers (unmyelinated; white ramus communicans) travel a short distance to the paravertebral sympathetic chain, and the postganglionic fibers (unmyelinated; gray ramus communicans) travel a relatively long distance to the effector organs. An exception to this rule is the adrenal medulla: playing, as it were, the role of a sympathetic chain ganglion, it receives long preganglionic fibers and then, instead of giving off postganglionic fibers, secretes epinephrine into the bloodstream. The parasympathetic preganglionic fibers are long; they project to ganglia near the effector organs, which, in turn, give off short postganglionic processes.

Neurotransmitters. Acetylcholine is the neurotransmitter in the sympathetic and parasympathetic ganglia. The neurotransmitters of the postganglionic fibers are norepineprhrine (sympathetic) and acetylcholine (parasympathetic). Neuromodulators include neuropeptides (substance P, somatostatin, vasoactive intestinal



lated peptide, neuropeptide Y, galanin, oxytocin, enkephalins) and nitric oxide.

141

The hypothalamus lies in the anterior portion of the diencephalon, below the thalamus and above the pituitary gland. It forms part of the wall and floor of the third ventricle. Among its anatomical components are the preoptic area, infundibulum, tuber cinereum, and mamillary bodies. It is responsible for the control and integration of endocrine function, thermoregulation (p. 152), food intake (p. 154), thirst, cardiovascular function (p. 148), respiration (p. 150), sexual function (p. 156), behavior and memory (p. 122 ff), and the sleep-wake rhythm (p. 112). Under the influence of changes in the external and internal environment, and the emotional state of the individual, the hypothalamus controls the activity of the ANS through its neural and humoral outflow.

Neuroendocrine Control

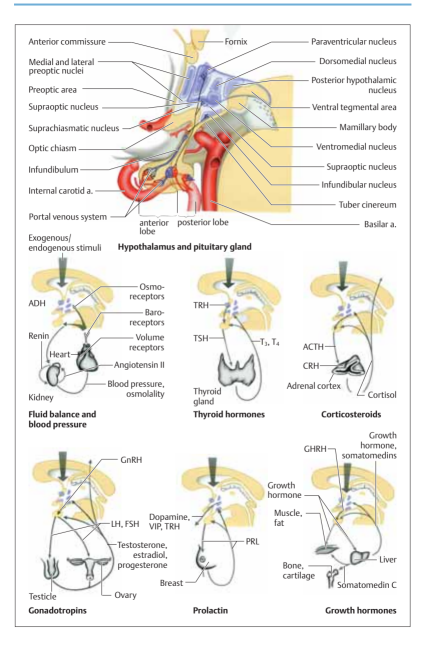
(Table 15, p. 367)

The neuroendocrine control circuits of the hypothalamic-pituitary axis regulate the plasma concentration of numerous hormones.

Adenohypophysis (anterior lobe of pituitary gland). Various regulatory hormones (releasing and inhibiting hormones) are secreted by hypothalamic neurons into a local vascular network, through which they reach the adenohypophysis to regulate the secretion of pituitary hormones into the systemic circulation. Among the pituitary hormones, the glandotropic hormones (TSH, ACTH, FSH, LH) induce the release of further hormones (effector hormones) from the endocrine glands, which, in turn, affect the function of the end organs, while the aglandotropic pituitary hormones (growth hormone, prolactin) themselves exert a direct effect on the end organs. Finally, the plasma concentration of the corresponding effector hormones and aglandotropic pituitary hormones affects the hypothalamic secretion of regulatory hormones in a negative feedback circuit (closed regulatory loop).

Neurohypophysis (posterior lobe of pituitary gland). A subset of hypothalamic neurons projects axons to the neurohypophysis. The bulblike endings of these axons store oxytocin and antidiuretic hormone and secrete them directly into the bloodstream (neurosecretion). These hormones act directly on their effector organs. The ensuing effects are sensed by the hypothalamus, thus closing the regulatory loop.

Hypothalamus



Limbic System

The limbic system consists of a number of separate structures with complex interconnections. Its function is only partly understood, but it is clear that it plays an important role in memory, emotion, and behavior.

Structure

The limbic system consists of inner and outer portions, both of which resemble a ring (Latin *limbus*). The outer portion extends from rostral structures (the septal and preoptic areas) in a craniocaudal arch (cingulate gyrus) to the temporal lobe, all the way to the temporal pole (hippocampus, entorhinal cortex). The inner portion extends from the hypothalamus and mamillary body via the fornix to the dentate gyrus, hippocampus, and amygdala.

Nerve pathways

The neuroanatomical loop, hippocampus \Rightarrow fornix \Rightarrow mammillary body \Rightarrow mammillothalamic tract \Rightarrow anterior thalamic nucleus \Rightarrow cingulate gyrus \Rightarrow cingulum \Rightarrow hippocampus, is called the Papez circuit. Numerous fiber tracts, many of them bilateral, connect the limbic system to the thalamus, cortex, olfactory bulb (p. 76), and brain stem. The medial forebrain bundle links the septal and preoptic areas with the hypothalamus and midbrain. Fibers from the amygdala pass in the stria terminalis, which occupies the groove between the caudate nucleus and the thalamus, to the septal area and hypothalamus. Short fibers from the amygdala also project to the hippocampus. The anterior commissure connects the two amygdalae, and the commissure of the fornix connects the two hippocampi.

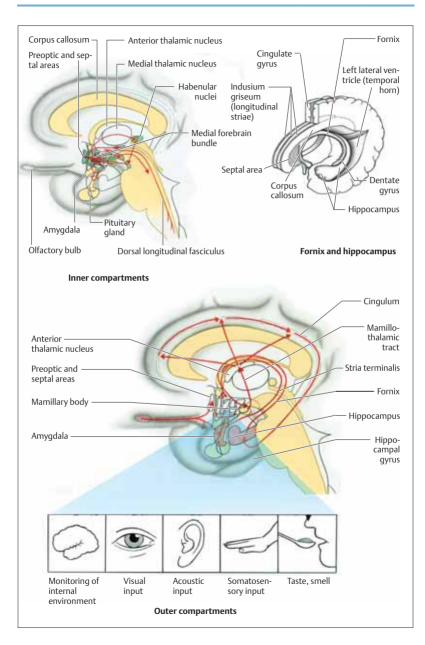
Functions of the limbic system

(Table 16, p. 368)

The limbic system controls emotional processes, such as those involved in anger, motivation, joy, sexuality, sleep, hunger, thirst, fear, aggression, and happiness. These processes are closely linked with cognition and memory (p. 134). The amygdala plays a key role in these events (emotional memory), integrating new, incoming information with the stored contents of memory. This integration determines the neural output of the limbic system, which affects the individual's physical state and behavioral responses.

ANS, Peripheral Portion (p. 146 ff)

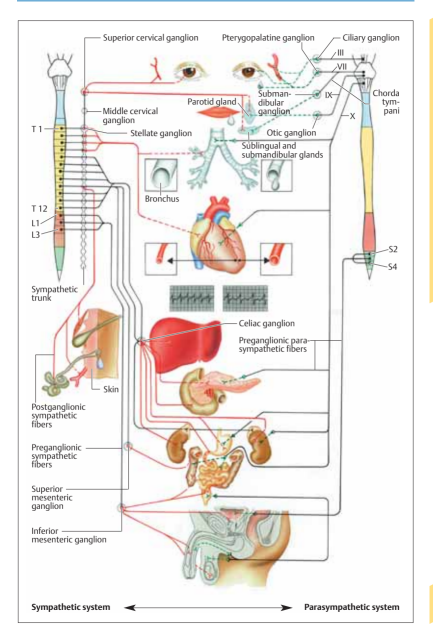
The peripheral portion of the ANS (p. 140) subserves a number of autonomic reflexes (p. 110). Nociceptive, mechanical, and chemical stimuli interact with their respective receptors to induce the generation of afferent impulses, which then travel to the spinal autonomic neurons. whose efferent output, as described in previous sections, controls the function of the heart, smooth muscles, and glands. The activity of the spinal sympathetic neurons is subject to supraspinal regulation by the autonomic centers of the brain. Most organs of the body receive both sympathetic and parasympathetic innervation. This double innervation enables synergistic coordination of multiple organ systems (e.g., acceleration of breathing and blood flow during physical exercise).



145

Sympathetic End Effect ¹ (Receptor Type ²)	Effector Organ	Parasympathetic End Effect ¹
Contraction ⇔ mydriasis (α ₁) Contraction ⇒ lid elevation — — —	Eye Dilator pupillae m. Tarsal m. Sphincter pupillae m. Ciliary m. Lacrimal gland	— Constriction ⇒ miosis (light response) Contraction ⇒ accommoda- tion (near response) Secretion
Light mucous secretion (α_1)	Salivary glands	Heavy serous secretion
Relaxation (β_2) Secretion: $\cup (\alpha_1), \cup (\beta_2)$ \cup Heart rate (β_1) \cup Contractility (β_1)	Thoracic organs Bronchial smooth muscle Bronchial glands Sinoatrial node Myocardium	Constriction D Secretion U Heart rate Slightly U contractility
Glycogenolysis, gluconeogenesis (α_1 , β_2) Dilatation (β_2) U Motility (α_2 , β_2) Contraction (α_1) U Insulin secretion (α_2) Renin secretion (α_2) Contraction (α_1) Relaxation (β_2) Contraction (pregnancy, α_1) Ejaculation (α_1)	Abdominal organs Liver Gallbladder, biliary tract Intestine Sphincters Pancreas Kidney Adrenal medulla Urinary sphincter Urinary detrusor Utreus Male genitalia	 Contraction Motility Relaxation Relaxation Contraction Varies (cycle-dependent) Erection
Secretion: Ω generalized (cholinergic), localized ⁴ Contraction (α_1)	Skin Sweat glands Arrector muscles of hair	-
Vasoconstriction (α_1, α_2) Vasoconstriction (α_1) , vasodilatation (β_2) Vasoconstriction (α_1) Vasoconstriction (α_1, α_2) , Vasodilatation (β_2) Vasoconstriction (α_1) Vasoconstriction (α_1, α_2)	Blood vessels Cutaneous arteries Arteries of skeletal muscle Cerebral arteries Coronary arteries Abdominal arteries Veins	Vasodilatation Vasodilatation Vasodilatation Slight vasoconstriction –

1 The action of the respective organ is listed in this column. **2** These are mainly membrane receptors for epinephrine and norepinephrine (adrenoceptors). Norepinephrine mainly acts on α and β_1 receptors; epinephrine acts on all types of adrenoceptors. *Sympathomimetic* \simeq increase sympathetic nervous activity (adrenoceptor agonist); *sympatholytic* \simeq reduces sympathetic nervous activity (adrenoceptor antagonist, receptor blocker); *parasympathomimetic* \simeq muscarinic receptor agonist; *indirect parasympathomimetic* \simeq blocks acetylcholinesterase; *parasympatholytic* (anticholinergic) \simeq muscarinic receptor antagonist; *antiparasympathotonic* \simeq botulinum toxin. **3** Preganglionic sympathetic fibers; transmitter acetylcholine. "Palms of hands (adrenergic sweating).



Heart and Circulation

Sympathetic efferent impulses cause arterial and venous vasoconstriction, acceleration of the heart rate, activation of the renin-angiotensinaldosterone system, and secretion of epinephrine/norepinephrine from the adrenal medulla. Blood pressure and blood volume rise. and blood is redistributed away from the vascular beds of skin (pallor), intestinal organs, and kidneys in favor of the heart and brain. Conversely, parasympathetic efferent impulses cause vasodilation and a decrease in heart rate. The normal arterial blood pressure is no higher than 140 mmHg systolic and 90 mmHg diastolic. Afferent connections. Baroreceptors (pressure sensors) are located between the media and adventitia of the arterial wall in the carotid sinus (innervated by CN IX), the aortic arch (X), and the brachiocephalic trunk (X). The impulses they generate are conveyed to the nucleus of the solitary tract (NST) in the dorsolateral medulla; the polysynaptic relay proceeds via interneurons to the anterolateral portion of the caudal medulla (CM), which in turn projects to inhibitory neurons in the anterolateral portion of the rostral medulla (RM). Other fibers from the NST project to the nucleus ambiguus (NA). Other baroreceptors, found in the atria and venae cavae near their entrance to the heart, sense the volume status of the vascular system and generate impulses that travel by way of CN X to the NST and hypothalamus. Cerebral ischemia (12 CO2 in extracellular fluid and CSF; p. 162) leads to increased sympathetic activity. Mechanical, nociceptive, metabolic, and respiratory influences also affect the medullary and hypothalamic centers that regulate the circulatory system.

Efferent connections. Sympathetic impulses travel from the (inhibitory) RM to the interomediolateral cell column of the spinal cord, which projects to the adrenal medulla and, through a relay in the sympathetic ganglia, to the heart, blood vessels, and kidney. The parasympathetic outflow of the NST is relayed in the nucleus ambiguus, by way of CN X, to the heart and blood vessels.

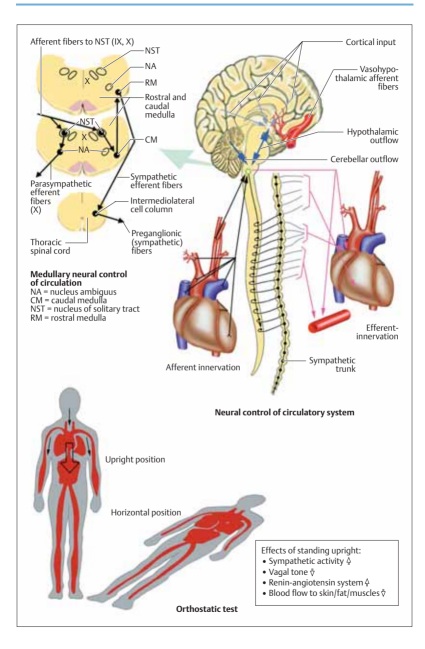
Central nervous regulation. The sympathetic and parasympathetic innervation of the circulatory system act synergistically. Both are ultimately controlled by the hypothalamus, which projects not only to the intermediolateral cell column (sympathetic), but also to the NST and nucleus ambiguus (parasympathetic). Afferent impulses from the skeletal muscles, baroreceptors, and vestibular organs reach the fastigial nucleus of the cerebellum, which has an excitatory projection to the NST and an inhibitory projection to the RM. Cortical projections to circulatory control centers enable the cardiovascular system to function as needed in the course of voluntary, planned movements.

Syndromes (Table 17, p. 369)

Neurogenic arrhythmias may be of S11praventricular or ventricular origin and are commonly associated with subarachnoid and intracerebral hemorrhage, head trauma. ischemic stroke, multiple sclerosis, epileptic seizures, brain tumors, carotid sinus syndrome (cardioinhibitory type), glossopharyngeal neuralgia and hereditary QT syndrome. They also sometimes occur in the immediate postoperative period after major neurosurgical procedures.

Neurogenic ECG abnormalities (ST depression or elevation, T-wave inversion) can occur in the setting of cerebral hemorrhage or infarction but are often difficult to distinguish from changes due to myocardial ischemia.

Hemodynamic abnormalities. Hypertension: Cerebral hemorrhage, Cushing reflex (accompanied by bradycardia) in response to elevated ICP, porphyria, Wernicke encephalopathy (accompanied by arrhythmia), and posterior fossa tumors. Hypotension: Head injuries, spinal lesions (syringomyelia, trauma, myelitis, funicular myelosis), multisystem atrophy, progressive supranuclear palsy, Parkinson disease, peripheral neuropathies (e.g., in diabetes mellitus, amyloidosis, Guillain-Barré syndrome, or renal failure). Neurocardiogenic syncope (vasovagal syncope) is due to pooling of venous blood in the arms and legs. Underfilling of the left ventricle activates baroreceptors, which, in turn, project via CN X to the NST. The ensuing increase of parasympathetic outflow, if large enough, sets off a "neurocardiogenic cascade" that leads to presyncope and syncope (p. 200). The characteristic features include decreased sympathetic activity, increased epinephrine secretion from the adrenal medulla, and decreased norepinephrine secretion, resulting in vasodilatation. Hyperexcitability of the vagus nerve (bradycardia) is also seen.



Respiration ensures an adequate oxygen supply for the body's tissues and maintains acid-base hemostasis.

Respiratory movements. Inspiration can be achieved by contraction of the diaphragm (*diaphragmatic respiration*) or of the intercostal muscles (*costal respiration*). The *auxiliary respiratory muscles* of the shoulder girdle further enlarge the chest cavity, if required, for deep breathing. Expiration is largely passive. The muscles of the abdominal wall and the latisimus dorsi muscle serve as *auxiliary expiratory muscles*. Other muscles (genioglossus, pharyngeal constrictor, and laryngeal muscles) keep the upper airway open during respiration.

Afferent connections. Chemoreceptors responding to arterial pH and O2 and CO2 concentration are found in the carotid glomus (innervated by CN IX), aortic arch (X), and para-aortic bodies (X). Impulses arising in these chemoreceptors, and in mechanoreceptors in the respiratory muscles (sensory afferent connections \Rightarrow phrenic nerve, intercostal nerves 2-12, CN IX) and in the lungs (bronchodilatation ⇒ pulmonary plexus, sympathetic nerve T1-T4), travel to the dorsal respiratory group of nuclei (DRG) anterior to the nucleus of the solitary tract, from which they are relayed, via interneurons, to the ventral respiratory group (VRG) in the medulla. The DRG also receives afferent input from the cardiovascular system, which is thus able to influence respiratory function. Changes in the pH and CO₂ concentration of the extracellular fluid (ECF) and cerebrospinal fluid (CSF) are also directly sensed by medullary chemoreceptors. Respiration is further influenced by a wide variety of other phenomena, e.g., cold, heat, hormones, reflexes (sneezing, coughing, yawning, swallowing), sleep, mental state (anxiety, fear), speaking, singing, laughing, muscle activity (physical work, sports), sexual activity, and body temperature.

Efferent connections. The laryngopharyngeal muscles and bronchoconstrictors are innervated by CN X. The phrenic nerve (diaphragm, C3–C5) and motor branches of the spinal nerves (intercostal nerves $2-9/T2-T11 \Rightarrow$ intercostal muscles; intercostal nerves $6-12/T6-T12 \Rightarrow$ abdominal wall muscles) supply the auxiliary respiratory muscles.

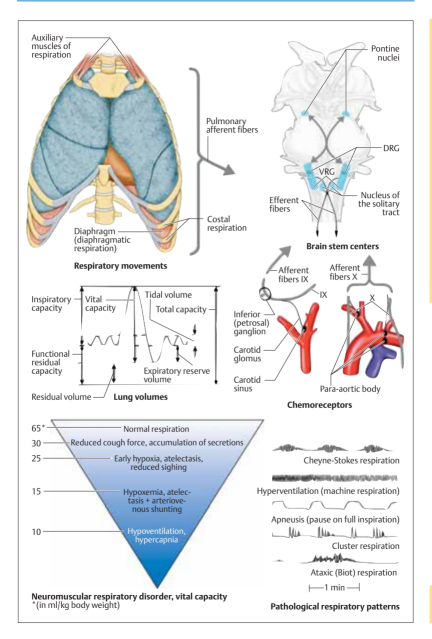
Respiratory rhythm. Rhythmic breathing is achieved by oscillating, alternately inhibitory

and excitatory *neural control circuits* within the VRG. As inspiration progresses, the inspiratory neuron groups of the VRG are progressively inhibited (Hering–Breuer reflex), while the expiratory neuron groups are excited. The respiratory rhythm is influenced by afferent input from chemoreceptors reflecting changes in the composition of the blood (decreases in pH and O_2 concentration, rise in CO₂ concentration \Rightarrow deeper respiratory rate) or of the ECF and CSF (decrease in pH, rise in CO₂ concentration \Rightarrow deeper respiration, Ω respiratory rate). The VRG is influenced by pontine nuclei.

Syndromes (Table 18, p. 370)

Pathological breathing patterns may be due to metabolic, toxic, or mechanical factors (obstructive sleep apnea) or to a lesion of the nervous system (p. 118). Morning headaches, fatigue, daytime somnolence, and impaired concentration may reflect a (nocturnal) breathing disorder. Neurogenic or myogenic breathing disorders often come to medical attention because of coughing attacks or food "going down the wrong pipe." Neurological diseases are often complicated by respiratory dysfunction. The respiratory parameters (respiratory drive, coughing force, blood gases, vital capacity) should be carefully monitored over time so that intubation and/or tracheostomy for artificial ventilation can be performed as necessary.

Respiration



The temperature of the body is a function of heat absorption, heat production, and heat elimination. Core body temperature normally fluctuates from approximately 36 °C in the morning to 37.5 °C in the late afternoon. It can be influenced by the menstrual cycle, pregnancy, and other hormonal factors, as well as by eating behavior and digestion, and it varies with age.

Neural control. The thermoregulatory center lies in the preoptic and anterior region of the hypothalamus (p. 142). Heat is eliminated through the skin (heat radiation, convection, sweating \Rightarrow cooling by evaporation), respiration (evaporation), and blood circulation (heat transport from the interior to the surface, cutaneous blood flow). Heat is produced by metabolic processes (under the influence of thyroid hormones) and by muscle contraction (shivering, voluntary movement). Thermoreceptors in the skin, spinal cord, medulla, and midbrain generate afferent impulses that travel to the hypothalamic control center; the cutaneous receptors project to the hypothalamus by way of the spinothalamic tract. Thermoreceptors are also present in the hypothalamus itself. The hypothalamus makes extensive connections with other regions of the brain. Its major efferent pathways relating to thermoregulation (vasomotor and sudoriparous pathways) pass by way of the ipsilateral lateral funiculus of the spinal cord to the spinal sympathetic nuclei (thoracolumbar system). These then give rise to fibers that travel by way of the ventral roots to the sympathetic chain ganglia; postganglionic fibers travel with the peripheral nerves to the skin. Sudoriparous fibers are found only in the ventral roots of T2/3 to L2/3, yet they innervate the skin of the entire body; thus, their distribution is not the same as the dermatomal distribution of sensation. Sudoriparous fibers to the head travel along the internal and external carotid arteries and then join branches of the trigeminal nerve to arrive at the skin. The neurotransmitter for the sympathetic innervation of the sweat glands is acetylcholine.

Disturbances of body temperature. *Central hyperthermia* is an elevation of body temperature due to impaired thermoregulation by the central nervous system. Its mechanism may involve either excessive heat production, excessive heat absorption (e.g., in a hot environment), or inadequate heat elimination. *Fever* may be defined as an oral temperature greater than 37.2 °C in the morning or 37.7 °C in the afternoon (rectal

temperature 0.6 °C higher). Its cause is generally not an impairment of thermoregulation, but rather a change in the set point for temperature established by the hypothalamic thermoregulatory center. Such a change can be brought about by circulating pyrogenic cytokines (e.g., interleukin-1, tumor necrosis factor, interferon- α) that exert an effect on hypothalamic function by interacting with the circumventricular organs (p. 140). *Hypothermia* is defined as a core temperature below 35 °C.

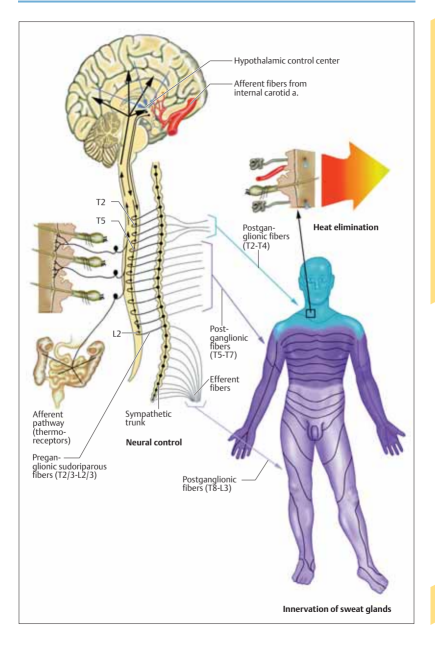
Syndromes

Disturbances of thermoregulatory sweating. *Examination:* Useful tests include palpation of the skin to appreciate its moisture and temperature, the quantitative sudomotor axon reflex test (QSART), the sympathetic skin response (SSR), iodine-starch test (Minor test), and the ninhydrin test.

Generalized anhidrosis (which confers a risk of hyperthermia) may be idiopathic or may be due to lesions in the hypothalamus or in the spinal cord above T3/4. Monoradicular lesions or cervical or lumbosacral polyradicular lesions do not impair sweating. Lesions of the sympathetic trunk cause segmental anhidrosis. *Plexus lesions* and *isolated or combined neuropathies* produce anhidrosis in the area of a sensory deficit. Lesions from the level of the stellate ganglion upward cause anhidrosis as a component of Horner syndrome. Sweating of the palms and soles is not influenced by thermoregulatory mechanisms but rather by the emotional state (fear, nervousness).

Central hyperthermia may be due to hypothalamic lesions (infarction, hemorrhage, tumor, encephalitis, neurosarcoidosis, trauma), intoxications (anticholinergic agents, salicy-lates, amphetamines, cocaine), acute spinal cord transection above T3/4, delirium, catatonia, malignant neuroleptic syndrome, malignant hyperthermia, dehydration, heat stroke, and generalized tetanus.

Fever. The symptoms include malaise, shivering, feeling cold, chills, nausea, vomiting, and somnolence. The heart rate and blood pressure rise, thermoregulatory sweating diminishes, and the peripheral blood volume is redistributed to the core of the body. Simple febrile convulsions in children under 5 years of age generally do not lead to epilepsy or other neurological complications.



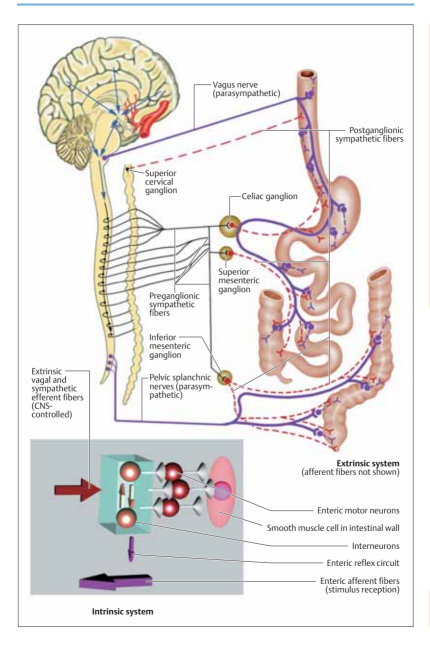
The uptake, transport, storage, and digestion of food, the absorption of nutrients, and the elimination of waste matter are under the influence of both the extrinsic autonomic nervous system and the intrinsic autonomic nervous system of the intestine.

The extrinsic system modulates the function of the intrinsic enteric system in coordination with the function of other organs of the body. Brain stem nuclei subserve the gastrointestinal and enterocolic reflexes, while cortical, limbic (hvpothalamus, amygdala) and cerebellar centers (fastigial nucleus) mediate the perception of satiety, the enteral response to hunger and odors, and emotional influences on alimentary function. The parasympathetic innervation (neurotransmitter: acetylcholine) of the esophagus, stomach, small intestine, and proximal portion of the large intestine is through the vagus nerve, while that of the distal portion of the large intestine and anal sphincter is derived from segments S2-S4. Parasympathetic activity stimulates intestinal motility (peristalsis) and glandular secretion. Sympathetic innervation (transmitter: norepinephrine) is from the superior cervical ganglion to the upper esophagus, from the celiac ganglion to the lower esophagus and stomach, and from the superior and inferior mesenteric ganglia to the colon. Sympathetic activity inhibits peristalsis, lowers intestinal blood flow, and constricts the sphincters of the gastrointestinal tract (lower esophageal sphincter, pylorus, inner anal sphincter).

The **intrinsic system (enteric system)** consists of the myenteric and submucous ganglionic plexuses (p. 141), which are independent neural networks of sensory and motor neurons and interneurons. The enteric autonomic nervous system receives chemical, nociceptive, and mechanical stimuli, processes this neural input, and produces efferent impulses affecting gastrointestinal glandular secretion and smooth-muscle contraction.

Syndromes (Table 19, p. 370)

Neurological diseases most commonly affect gastrointestinal function by impairing motility, less commonly by impairing resorptive and secretory processes. The differential diagnosis must include gastrointestinal dysfunction of nonneurological (often obstructive) origin. Specialized diagnostic testing is usually indicated.



Bladder Function

The urinary bladder stores (continence) and voids (micturition) the urine produced by the kidneys. Parasympathetic fibers arising in segments S2-S4 (sacral micturition center, detrusor nucleus) and traveling through the pelvic plexus activate the detrusor muscle of the bladder. Sympathetic fibers arising from segments T10-L2 and traveling through the hypogastric plexus inhibit the detrusor (β-adrenergic receptors) and stimulate the vesical neck (trigone, internal sphincter; α -adrenergic receptors). Somatic motor impulses arising from segments S2-S4 (Onuf's nucleus) travel through the pudendal nerve to the external sphincter and the pelvic floor muscles. Somatosensory fibers from the bladder travel along the hypogastric and pelvic nerves to spinal levels T10-L2 and S2-S4, conveying information about the state of bladder stretch (overdistention is painful). The central nervous system (frontal lobes, basal ganglia) subserves the voluntary inhibition of detrusor contraction. The pontine micturition center, which triggers the act of micturition, is under the influence of afferent impulses relating to the state of bladder stretch; its output passes to somatic motor neurons in the spinal cord that synergistically innervate the detrusor and external sphincter muscles.

Continence. An intact bladder closure mechanism is essential for normal filling (up to 500 ml). Closure involves contraction of the vesical neck and external sphincter urethrae and pelvic floor muscles, and relaxation of the detrussor muscle (dome of the bladder).

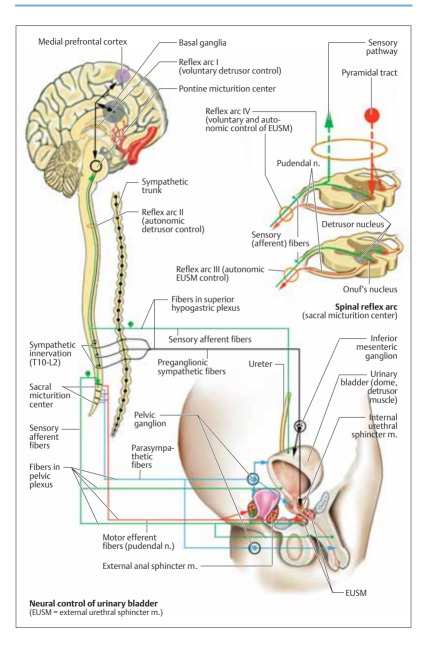
Micturition. Once a urine volume of 150–250 ml has accumulated, stretch receptors generate impulses that pass to the pontine micturition center and also produce the sensation of bladder distention. Micturition begins when the dome of the bladder is stimulated to contract while the vesical neck and pelvic floor muscles relax. Contraction of the muscles of the abdominal wall increases the intravesical pressure and facilitates micturition.

Neurogenic bladder dysfunction (Table 20, p. 371). Additional diagnostic tests are performed in collaboration with a urologist or gynecologist as deemed necessary on the basis of the case history and neurological findings.

Useful diagnostic aids include laboratory testing (urinalysis and renal function), ultrasound examination (kidney, bladder, pelvis), urodynamic testing, micturition cystourethrography, and neurophysiological studies (evoked potentials, urethroanal/bulbocavernosus reflex).

Sexual Function

The genital organs receive sympathetic (T11– L2), parasympathetic (S2–S4), somatic motor (Onuf's nucleus), and somatosensory innervation (S2–S4) and are under supraspinal control, mostly through hypothalamic projections to the spinal cord. Hormonal factors also play an important role (p. 142). Neurological disease often causes sexual dysfunction (erectile dysfunction, ejaculatory dysfunction) in combination with bladder dysfunction. Isolated sexual dysfunction is more often due to psychological factors (depression, anxiety), diabetes mellitus, endocrine disorders, and atherosclerosis.



The intracranial pressure (ICP) corresponds to the pressure that the contents of the skull exert on the dura mater.

Intracranial Hypertension

The normal intracranial pressure (ICP) is 60-120 mmH₂O, which corresponds to 5-15 mmHg. An ICP greater than 30 mmHg impairs cerebral blood flow; an ICP greater than 50 mmHg for more than 30 minutes is fatal; an ICP greater than 80 mmHg for any length of time can cause brain damage. Intracranial hypertension may be either acute (developing in hours to days) or chronic (lasting for weeks or months). Its manifestations are progressively more severe as the ICP rises, but are not specific; thus, the diagnosis cannot be made from the signs of intracranial hypertension alone but requires either the demonstration of a causative lesion (e.g., subdural hematoma, encephalitis, brain tumor, hydrocephalus) or direct measurement of the intracranial pressure. Treatment is indicated when the ICP persistently exceeds 20 mmHg, when plateau waves are found, or when the pulse amplitude rises. Individual cases may manifest a variety of different signs of intracranial hypertension, either in slow or rapid alternation, or all at the same time. Compression of the brain stem by a space-occupying lesion (p. 118) has similar clinical manifestations; the diagnostic differentiation of brain stem compression from intracranial hypertension is critical. Lumbar puncture is contraindicated in cases of suspected or documented intracranial hypertension, as the resulting increase in the already high craniospinal pressure gradient may lead to brain herniation.

■ Clinical Features (Table 21, p. 371)

Headache due to intracranial hypertension ranges in intensity from mild to unbearable. Patients typically report a pressing, bifrontal headache that is most severe upon awakening in the morning or after naps in the daytime. It is exacerbated by lying flat, coughing, abdominal straining, or bending over, and ameliorated by sitting or standing. It may wake the patient from sleep. Often, both mild daytime headaches and more severe nighttime headaches are present.

Nausea due to intracranial hypertension is often independent of movement of the head or of

other abdominal complaints, and its intensity is not correlated with that of headache. It may be mild or severe.

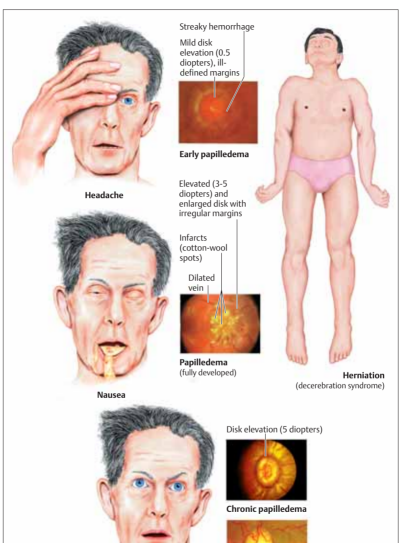
Projectile vomiting may occur without warning or after a brief sensation of nausea upon sitting up or moving the head. Initially, vomiting mainly occurs suddenly, in the morning (on an empty stomach).

Eye movements and vision. Compression of CN III or VI causes paresis of extraocular muscles or pupillary dilatation (p. 92). Papilledema often affects the eve on the side of the causative lesion first, and then gradually the other eye as well. The older the patient, the less likely that papilledema will occur as a sign of intracranial hypertension; its absence thus cannot be taken as ruling out intracranial hypertension. Early papilledema is characterized by hyperemia, blurred papillary margins, dilated veins, loss of venous pulsations (may be absent normally), and small hemorrhages around the papilla. Full-blown papilledema is characterized by disk elevation, engorged veins, tortuous vessels around the papilla, and streaky hemorrhages. If intracranial hypertension persists, chronic papilledema will develop in weeks or months, characterized by gravishwhite optic nerve atrophy and small vessel caliber. Acute papilledema generally does not affect the visual fields or visual acuity (unlike papillitis, which should be considered in the differential diagnosis); but physical exertion or head movement may cause transient amblyopic attacks lasting several seconds (foggy or blurred vision, or blindness). Chronic papilledema, on the other hand, can cause an impairment visual acuity, concentric visual field defects, and even blindness,

Gait disturbances. An unsteady, slow, hesitant, gait with small steps and swaying from sided to side is sometimes seen.

Behavioral changes. Impairment of memory, attention, concentration, and planning ability, confusion, slowed reactions, and changes in personal habits are often observed by relatives and friends.

Intracranial Pressure



159

Behavioral change



Optic nerve atrophy

Herniation syndromes (p. 162). Transtentorial herniation causes an ipsilateral oculomotor nerve palsy (ptosis, mydriasis, and secondary ophthal-moplegia), contralateral hemiplegia, and decerebration syndrome (p. 46). Downward herniation of the contents of the posterior fossa into the foramen magnum causes neck pain and stiffness, a head tilt, and shoulder paresthesiae. If medullary compression is also present, respiratory and circulatory disorders, cerebellar fits, and obstructive hydrocephalus may develop. Upward herniation of the contents of the posterior fossa across the tentorial notch causes a decerebration syndrome in which the ipsilateral pupil is initially constricted and later dilated.

Pseudotumor cerebri causes headache (holocephalic, bilateral frontal/occipital), visual disturbances of varying severity (enlarged blind spot, blurred vision, loss of vision, or diplopia due to abducens palsy), and bilateral papilledema. CT and MRI scans reveal the absence of an intracranial mass (normal ventricular size, thickening of optic nerve, "empty sella" = intrasellar expansion of the suprasellar cisterns, with or without sellar dilatation). CSF tests are normal except for an elevated opening pressure (>250 mmH₂O). The etiology of pseudotumor cerebri is multifactorial; it occurs most commonly in obese young women. Its differential diagnosis includes intracranial venous or venous sinus thrombosis, drug toxicity (high doses of vitamin A, tetracycline, NSAIDs), elevated CSF protein level (spinal tumor, Guillain-Barré syndrome), or endocrine changes (pregnancy, Addison disease, Cushing syndrome, hypothyroidism).

Intracranial Hypotension

Spontaneous drainage of cerebrospinal fluid by means of lumbar puncture is no longer possible when the CSF pressure is below 20 mmH₂O (patient is lying flat); when it is below 0 mmH₂O, air is sucked through the LP needle into the subarachnoid space and travels upward to the head, where air bubbles can be seen on a CT scan. For causes of low intracranial pressure see Table 22 (p. 372).

Symptoms

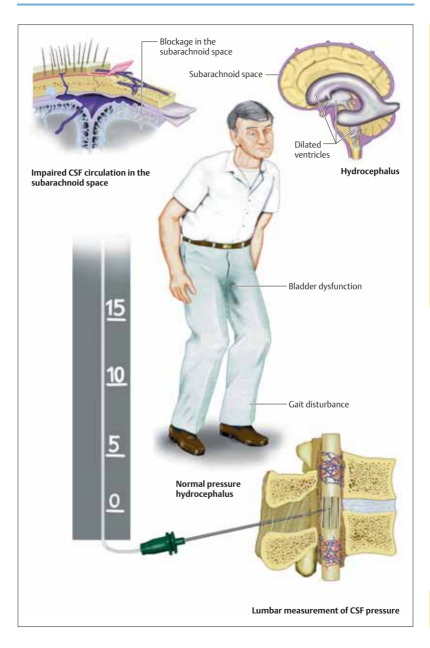
Severe headache (nuchal, occipital, or frontal) is provoked by sitting, standing, or walking, and subsides when the patient lies flat. It is exacerbated by abdominal straining, coughing, and the Valsalva maneuver. Other symptoms include nausea, vomiting, and dizziness. Unilateral or bilateral abducens palsy, tinnitus, ear pressure, or neck stiffness may also occur. Subdural fluid collections (hematoma due to rupture of the bridging veins, hygroma arising from local CSF collection due to rupture of the arachnoid membrane) are rare. Intracranial hypotension may be caused by a CSF leak; patients may report occasional loss of watery fluid from the nose or ear (traces visible on pillows).

Normal-pressure hydrocephalus (NPH) is a chronic form of communicating hydrocephalus that reflects an impairment of CSF circulation and resorption, sometimes in the aftermath of subarachnoid hemorrhage, head trauma, or chronic meningitis, but often without discernible cause. Symptoms develop over several weeks or months. Gait disturbances (gait apraxia, hydrocephalic astasia-abasia) begin as unsteadiness, difficulty climbing stairs, leg fatigue, a small-stepped gait, and frequent stumbling and falling, and then typically progress to an inability to stand, sit, or turn over in bed. The associated behavioral changes (p. 132 ff) are variable and may include impaired spatial orientation. reduced psychomotor drive (abulia), mild memory disturbances, or even dementia. Bladder dysfunction such as urge incontinence and polyuria develop as the condition progresses. Patients ultimately lose the perception of bladder distension and thus void uncontrollably.

Pathogenesis

ICP depends on the volume of nervous tissue, CSF, and blood inside the nondistensible cavity formed by the skull and vertebral canal. An increase in the volume of one of these three components must be offset by a compensatory mechanism (Monro-Kellie doctrine) such as an adjustment of the CSF or (venous) blood volume, expansion or the lumbosacral dural sheath, or deformation of the brain. When the capacity of such mechanisms is exhausted, the ICP decompensates. *Compliance* is defined as the first derivative of volume as a function of ICP, i.e., the ratio of a small, incremental change in

Intracranial Pressure



volume to the change in ICP that it produces. Compliance is thus an index of the ability to compensate for changes in volume. Decreased compliance leads to ICP decompensation. *Elastance* is the reciprocal of compliance, hence indicating the inability to compensate for changes in volume.

CSF volume. Hydrocephalus is defined as abnormal dilatation of the ventricular system. It occurs because of disturbances of CSF circulation and/or resorption. Hydrocephalus due to a blockage of the CSF pathway at some point within the ventricular system is called noncommunicating or obstructive hydrocephalus. Hydrocephalus due to impaired CSF resorption at the arachnoid villi is called communicating or malresorptive hydrocephalus. Acute hydrocephalus is characterized by ventricular dilatation with acute intracranial hypertension. Hydrocephalus with normal ICP and without progression of ventricular dilatation is called arrested, compensated or chronic hydrocephalus. So-called normal-pressure hydrocephalus occupies an intermediate position between these two conditions. External hydrocephalus is a dilatation of the subarachnoid space with no more than mild enlargement of the ventricles.

Cerebral Blood Flow

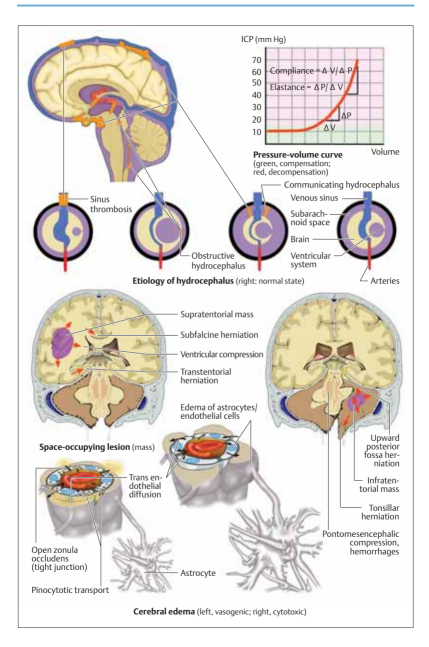
Cerebral blood flow (CBF) is a function of the cerebral perfusion pressure (CPP), which normally ranges from 70 to 100 mmHg, and the cerebral vascular resistance (CVR): CBF = CPP/CVR. CBF is maintained at a constant value of approximately 50 ml/100 g/min as long as the mean arterial pressure (MAP)¹ remains in the relatively wide range from 50 to 150 mmHg (cerebral autoregulation). When the patient is lying flat, CPP = MAP - ICP. A rapid rise in the systemic arterial pressure is followed by a slow, delayed rise in ICP; chronic arterial hypertension usually does not affect the ICP. On the other hand, any elevation of the venous pressure (normal central venous pressure: 40-120 mmH₂O) due to Valsalva maneuvers, hypervolemia, right heart failure, changes in body position, or obstruction of jugular venous drainage elevates the ICP by a comparable amount. Acidosis (defined as pH < 7.40), which may be due to hypoxia, ischemia, or hypoventilation, causes cerebral vasodilatation and increased cerebral blood volume, thereby elevating the ICP. Chronic obstructive pulmonary disease can elevate the ICP by this mechanism. On the other hand, *alkalosis* (e. g., due to hyperventilation) reduces the cerebral blood volume and thus also the ICP. CBF and ICP are elevated in fever and low in hyperthermia.

Intracranial Space-occupying Lesions

Extra-axial or intra-axial compression of brain tissue elevates the ICP, calling compensatory mechanisms into play: once these have been exhausted, mass displacement of brain tissue occurs, possibly resulting in herniation. The distribution of pressure within the cranial cavity is a function of the structure of the brain and the partitioning of the cavity by dural folds (p.6). Different herniation syndromes occur depending on the site and extent of the causative lesion: subfalcine herniation involves movement of the cingulate gyrus under the falx cerebri; transtentorial herniation involves movement of the medial portion of the temporal lobe across the tentorial notch: upward posterior fossa herniation involves movement of the brain stem and cerebellum across the tentorial notch: and downward posterior fossa herniation involves movement of the cerebellar tonsils across the foramen magnum.

In cerebral edema, accumulation of water and electrolytes in brain tissue causes an increase in brain volume. Vasogenic cerebral edema is due to increased capillary permeability and mainly affects white matter; it is caused by brain tumor, abscess, infarction, trauma, hemorrhage, and bacterial meningitis. Cytotoxic cerebral edema affects both white and gray matter and is due to fluid accumulation in all cells of the brain (neurons, glia, endothelium) because of hypoxia/ ischemia or acute hypotonic hyperhydration (water intoxication, dysequilibrium syndrome, inadequate ADH syndrome). Hydrocephalic (interstitial) brain edema is found in the walls of the cerebral ventricles and results from movement of fluid from the ventricles into the adiacent tissue in the setting of acute hydrocephalus.

¹The MAP can be estimated with the following formulas: MAP = [(systolic BP) + (2 × diastolic BP)] × ¹/₃ or, equivalently, MAP = diastolic BP + [(BP amplitude) × ¹/₃]



3 Neurological Syndromes

- Brain Disorders
- Spinal Disorders
- Peripheral Neuropathies
- Myopathies

A *stroke* is an acute focal or global impairment of brain function resulting from a pathological process (e.g. thrombus, embolus, vessel rupture) of the blood vessels. Its causes, in order of decreasing frequency, are *ischemia* (80%), spontaneous *intracerebral or intraventricular hemorrhage* (15%), and *subarachnoid hemorrhage* (5%). The signs and symptoms of stroke are usually not specific enough to enable identification of its etiology without further diagnostic studies. CT, MRI, cerebrovascular ultrasonography, ECG, and laboratory testing are usually needed.

Symptoms and Signs

The clinical manifestations of stroke persist, by definition, for more than 24 hours, and are often permanent, though partial recovery is common. The duration of symptoms and signs seems not to be correlated with the etiology of stroke.

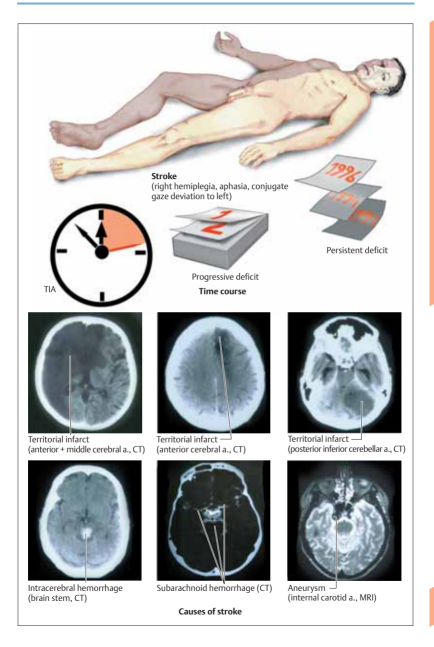
Ischemia. A transient ischemic attack (TIA) differs from a stroke (by definition) in that its symptoms and signs resolve completely within 24 hours. The vast majority of TIAs resolve within one hour, and only 5% last longer than 12 hours. Patients with crescendo TIAs (a rapid succession of TIAs) have a high risk of developing a (completed) stroke, which can cause neurological deficits that are either minor (minor stroke) or major (disabling stroke, major stroke). A stuttering, fluctuating, or progressive course of stroke development (*stroke in evolution*) is uncommon.

Hemorrhage. Nontraumatic intracerebral hemorrhages usually cause acute neurological deficits that persist thereafter. If deficits worsen after the initial hemorrhage, the cause is either recurrent hemorrhage or a complication of the initial hemorrhage (cerebral edema, electrolyte imbalance, or heart disorder).

Type of Deficit	Clinical Manifestations
Weakness (pp. 46 ff, 70)	Acute hemi-, mono-, or quadriparesis/quadriplegia (ca. 80–90%); loss of coordination and balance; hyperkinesia (during or after stroke), e.g. hemichorea, hemiballism, or (rarely) dystonia
Sensory loss (pp. 70 ff, 106)	Injury of postcentral cortex or subcortical area \Rightarrow distal sensory (often also motor) deficit in contralateral limbs. Paresthesiae and loss of stereognosis, graphesthesia, topesthesia, and acrognosis are prominent
Oculomotor and visual disturbances (pp. 70, 82 ff)	Conjugate horizontal eye movements, disjugate gaze, nystagmus, diplopia. Visual field defects (p. 82), transient monocular blindness (= amaurosis fugax)
Headache (p. 182)	May be caused by subarachnoid hemorrhage, temporal arteritis, venous sinus thrombo- sis, arterial dissection, cerebellar hemorrhage, massive intracerebral hemorrhage (rare)
Impairment of con- sciousness (pp. 116, 204)	TIA and stroke generally do not impair consciousness (exception: brainstem stroke, massive supratentorial stroke with bilateral cortical dysfunction)
Behavioral changes (p. 122 ff)	Aphasia, confusion (must be distinguished from aphasia), impairment of memory, neg- lect, impaired affect control (compulsive crying and/or laughing), apraxia. Mental changes, especially depression and anxiety disorders, are common after stroke
Dysarthria and dysphagia (pp. 102, 130)	Severe dysarthria is often accompanied by coughing, difficulty chewing, and dysphasia. <i>Pseudobulbar palsy</i> \Rightarrow loss of voluntary motor control (e.g., swallowing, speaking, tongue movement) with preservation of involuntary movements (e.g., yawning, coughing, laughing)
Dizziness (p. 58)	Cerebellum, brainstem (vertigo, nausea, nystagmus)
Epileptic seizures (p. 192 ff)	Simple partial, complex partial, or generalized tonic-clonic seizures may occur during or after a stroke
Respiratory disorders (p. 150)	<i>Hiccups</i> (singultus) often occur in stroke, particularly in lateral medullary infarction. Central hyperventilation is associated with a poor prognosis. Bihemispheric lesions may cause Cheyne–Stokes respiration (p. 118)

Minor complications of stroke: Mild unilateral arm paresis, moderate sensory loss, mild dysarthria; these patients can care for themselves. Mojor complications: Aphasia, spastic hemiplegia, and hemianopsia; these patients generally need nursing care.

Stroke



167

Stroke Syndromes: Carotid Artery Territory

Brachiocephalic Trunk

Brachiocephalic trunk occlusion by emboli from the aortic arch has the same clinical manifestations as internal carotid artery (ICA) occlusion. Patients with adequate collateral flow remain asymptomatic.

Common Carotid Artery (CCA)

CCA occlusion is very rare and, even when it occurs, is usually asymptomatic, because of an adequate collateral supply. When symptoms do occur, they are the same as those of ICA occlusion.

Internal Carotid Artery (ICA)

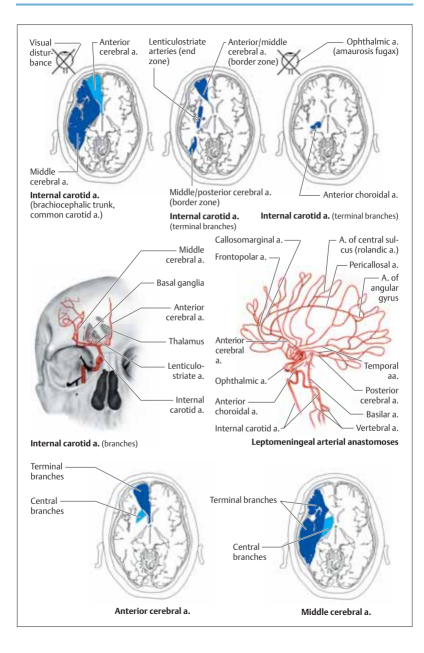
Territorial infarcts affect the middle cerebral artery (MCA) more often than the anterior cerebral artery (ACA). If the ICA is occluded and collateral flow via the circle of Willis is inadequate, extensive infarction occurs in the anterior twothirds of the hemisphere, including the basal ganglia. Symptoms include partial or total blindness in the ipsilateral eye, impairment of consciousness (p. 116), contralateral hemiplegia and hemisensory deficit, homonymous hemianopsia, conjugate gaze deviation to the side of the lesion, and partial Horner syndrome. ICA infarcts in the dominant hemisphere produce global aphasia. The occipital lobe can also be affected if the posterior cerebral artery (PCA) arises directly from the ICA (so-called fetal origin of the PCA). Border zone infarcts occur in distal vascular territories with inadequate collateral flow. They affect the "watershed" areas between the zones of distribution of the major cerebral arteries in the high parietal and frontal regions, as well as subcortical areas at the interface of the lenticulostriate and leptomeningeal arterial zones.

Ophthalmic artery. Occlusion leads to sudden blindness ("black curtain" phenomenon or centripetal shrinking of the visual field), which is often only temporary (*amaurosis fugax* = transient monocular blindness). Thorough diagnostic evaluation is needed, as the same clinical syndrome can be produced by other ophthalmological diseases (Table 22a, p. 372).

Anterior choroidal artery (AChA). Infarction in the AChA territory, depending on its precise location and extent, can produce contralateral motor, sensory, or mixed deficits, hemiataxia, homonymous quadrantanopsia (both upper and lower), memory impairment, aphasia, and hemineglect.

Anterior cerebral artery (ACA). Contralateral hemiparesis is usually more distal than proximal, and more prominent in the lower than in the upper limb (sometimes only in the lower limb). Infarction in the territory of the central branches of the ACA (A1 segment, recurrent artery of Heubner) produces brachiofacial hemiparesis, sometimes accompanied by dystonia. Bilateral ACA infarction (when the arteries of both sides share a common origin) and infarctions of the cortical branches of the ACA produce abulia (p. 120), Broca aphasia (dominant hemisphere), perseveration, grasp reflex, palmomental reflex, paratonic rigidity (gegenhalten), and urinary incontinence. Lesions in the superior and medial frontal gyri or the anterior portion of the cingulate gyrus cause bladder dysfunction. Disconnection syndromes due to lesions of the corpus callosum are characterized by ideomotor apraxia, dysgraphia, and tactile anomia of the left arm.

Middle cerebral artery (MCA). Main trunk (M1) occlusion produces contralateral hemiparesis or hemiplegia with a corresponding hemisensory deficit, homonymous hemianopsia, and global aphasia (dominant side) or contralateral hemineglect with limb apraxia (nondominant side). Occlusion of the posterior main branch produces homonymous hemianopsia or quadrantanopsia as well as Wernicke or global aphasia (dominant side) or apraxia and dyscalculia (nondominant side); central main branch occlusion produces contralateral brachiofacial weakness and sensory loss; anterior branch occlusion on the dominant side additionally produces Broca aphasia. Occlusion of peripheral branches produces monoparesis of the face, hand, or arm. Occlusions of the lenticulostriate arteries, depending on their precise location, produce (purely motor) hemiparesis/hemiplegia, or hemiparesis with ataxia (lacunar infarct, p. 172).



Stroke Syndromes: Vertebrobasilar Territory

Subclavian Artery

High-grade subclavian stenosis or occlusion proximal to the origin of the vertebral artery may cause a reversal of blood flow in the vertebral artery, which worsens with exertion of the ipsilateral arm (*subclavian steal*). Rapid arm fatigue and pain often result; less common are vertigo and other brain stem signs. The arterial blood pressure is measurably different in the two arms.

Vertebral Artery (VA)

VA occlusion produces variable combinations of symptoms and signs, including homonymous hemianopsia, dysarthria, dysphagia, unilateral or bilateral limb paralysis with or without sensory deficit, ataxia, drop attacks (due to medullary ischemia), and impairment of consciousness. Unilateral VA occlusion (e.g., due to dissection) can lead to infarction in the territory of the posterior inferior cerebellar artery.

Cerebellar Arteries

Large cerebellar infarcts can cause brain stem compression and hydrocephalus.

Posterior inferior cerebellar artery (PICA). Dorsolateral medullary infarction produces (usually incomplete) *Wallenberg syndrome* (p. 361). Often, only branches to the cerebellum are affected (⇔ vertigo, headache, ataxia, nystagmus, lateropulsion).

Anterior inferior cerebral artery (AICA). AICA occlusion is rare. It produces ipsilateral hearing loss, Horner syndrome, limb ataxia, and dissociated facial sensory loss, as well as contralateral dissociated sensory loss on the trunk and limbs (mainly the upper limbs) and nystagmus.

Superior cerebellar artery (SCA). SCA occlusion can produce ipsilateral Horner syndrome, limb ataxia, dysdiadochokinesia, and CN VI and VII palsy, as well as contralateral hypesthesia and hypalgesia.

Basilar Artery (BA)

Basilar artery occlusion. Thrombotic occlusion of the BA may be heralded several days in advance by nonspecific symptoms (unsteadiness, dysarthria, headache, mental changes). BA occlusion causes impairment of consciousness (ranging from somnolence to coma), mental syndromes (hallucinations, confabulation, psychoses), quadriparesis, and oculomotor disorders (diplopia, vertical or horizontal gaze palsy). Apical BA occlusion (p. 359) is caused by cardiac or arterial emboli. *Pontine infarction* sparing the posterior portion of the pons (tegmentum) produces quadriplegia and mutism with preservation of sensory function and vertical eye movements (*locked-in syndrome*, pp. 120, 359).

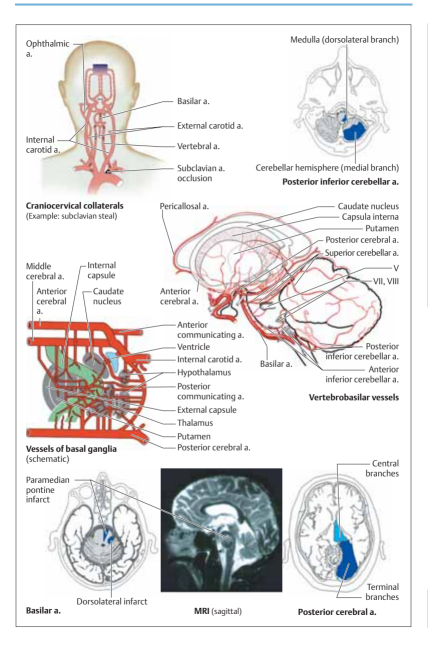
Paramedian infarction in the BA territory usually affects the pons (pp. 72, 359 ff).

Dorsolateral infarction affects the cerebellum, with a corresponding clinical picture. Occlusion of the labyrinthine artery (a branch of the AICA) produces rotatory vertigo, nausea, vomiting and nystagmus.

Posterior Cerebral Artery (PCA)

PCA occlusion is rare and produces symptoms and signs similar to those of MCA infarction. Unilateral occlusion of a cortical branch produces homonymous hemianopsia with sparing of the macula (supplied by the MCA), while bilateral occlusion produces cortical blindness and, occasionally, Anton syndrome (p. 132). Central branch occlusion leads to thalamic infarction (p. 106; Dejerine-Roussy syndrome), resulting in transient contralateral hemiparesis, spontaneous pain ("thalamic pain"), sensory deficits. ataxia, abasia, choreoathetosis, "thalamic hand" (flexion of the metacarpophalangeal joints with hyperextension of the interphalangeal joints), and homonymous hemianopsia. If branches to the midbrain are affected, an ipsilateral CN III palsy results, accompanied by variable contralateral deficits including hemiparesis/hemiplegia, (rubral) tremor, ataxia, and nystagmus. Isolated hemihypesthesia is associated with thalamic lacunar infarction.

Stroke: Ischemia



Risk Factors

The risk of stroke increases with age and is higher in men than in women at any age. Major risk factors include arterial hypertension (>140 mmHg systolic, >90 mmHg diastolic), diabetes mellitus, heart disease, cigarette smoking, hyperlipoproteinemia (total cholesterol >5.0 mmol/l, LDL >3 mmol/l, HDL <0.9-1.2 mmol/l), elevated plasma fibrinogen, and obesity. Symptomatic or asymptomatic carotid artery stenosis, elevated plasma homocysteine levels, erythrocytosis, anti-phospholipid antibodies, alcohol abuse (≥ 60 g of alcohol ≈ 75 cl of wine per day in men, ≥ 40 g in women). Drug abuse (amphetamines, heroin, cocaine), a sedentary lifestyle, and low socioeconomic status (unemployment, poverty) also increase the risk of stroke.

Causes

Embolism (ca. 70%) is the most common cause of stroke. Emboli arise from local atheromatous lesions (atheromatous thromboembolism) on the walls of large arteries (macroangiopathy) of the brain or heart (cardiac embolism in atrial fibrillation, valvular heart disease, ventricular thrombus, and myxoma).

Thrombosis (ca. 25%). Occlusion of a small endartery (microangiopathy, small vessel disease) causes lacunar infarction. The cause is hyaline (lipohyalinosis) or proximal sclerosis of penetrating arteries (lenticulostriate, thalamoperforating or pontine arteries, central branches). Causal factors include hypertension, diabetes, and bloodbrain barrier disruption leading to deposition of plasma proteins in the arterial wall. Microangiopathy-related hemodynamic changes sometimes cause hemodynamic infarction.

Rare causes (ca. 5%) include hematological diseases (e.g., coagulopathy, abnormal blood viscosity, anemia, leukemia) and arterial processes (dissection, vasculitis, migraine, fibromuscular dysplasia, moyamoya, vasospasm, amyloid angiopathy, and CADASIL = cerebral autosomal dominant arteriopathy with subcortical infarcts and leukoencephalopathy).

Infarct Types

Lacunar infarcts ("small deep infarcts"). Lacunes are small (\leq 1.5 cm in diameter), round or oval

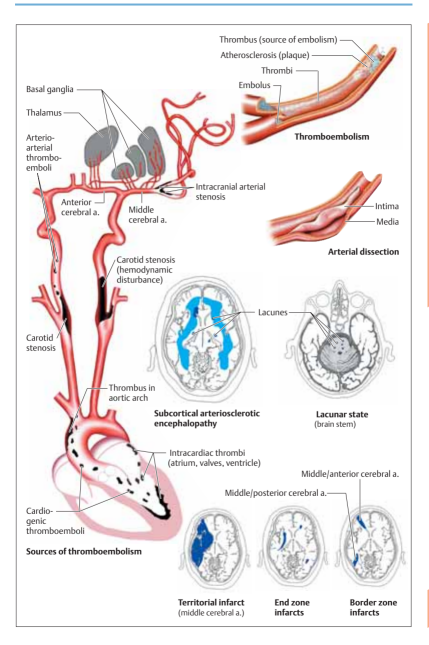
infarcts in the subcortical periventricular region or brain stem. Classic lacunar syndromes include purely motor hemiparesis (internal capsule, corona radiata, pons), contralateral purely sensory deficit (thalamus, internal capsule), ataxic hemiparesis (internal capsule, corona radiata, pons), and dysarthria with clumsiness of one hand (= clumsy hand-dysarthria syndrome; internal capsule, pons). The presence of multiple supratentorial and infratentorial lacunes is termed the lacunar state ("état lacunaire") and is clinically characterized by pseudobulbar palsy (p. 367), small-step gait ("marche à petit pas"), urinary incontinence, and affective disorders (compulsive crying). For leukoaraiosis, see p. 298.

Territorial infarcts are those limited to the distribution of the ACA, MCA, or PCA. With the exception of striatocapsular infarcts (internal capsule, basal ganglia), these infarcts are predominantly cortical. Embolic territorial infarcts often undergo secondary hemorrhage ("hemorrhagic conversion").

End zone infarcts. *Low-flow infarction* in the subcortical white matter is due to extracranial high-grade vessel stenosis and/or inadequate collateral flow.

Border zone infarcts (p. 168) also result from hemodynamic disturbances due to microangiopathy. They are found at the interface ("watershed") between adjacent vascular territories, and can be either anterior (MCA–ACA \Rightarrow contralateral hemiparesis and hemisensory deficit, mainly in the lower limb and sparing the face, with or without aphasia) or posterior (MCA– PCA \Rightarrow contralateral hemianopsia and cortical sensory deficit, with or without aphasia).

Global cerebral hypoxia/ischemia. The causes include cardiac arrest with delayed resuscitation, hemorrhagic shock, suffocation, and carbon monoxide poisoning. Global cerebral hypoxia/ischemia causes bilateral necrosis of brain tissue, particularly in the basal ganglia and white matter.



Stroke Pathophysiology

Hemodynamic insufficiency. Cerebrovascular autoregulation is normally able to maintain a relatively constant cerebral blood flow (CBF) of 50-60 ml/100 g brain tissue/min as long as the mean arterial pressure (MAP) remains within the range of 50-150 mmHg (p. 162). The regional cerebral blood flow (rCBF) is finely adjusted according to local metabolic requirements (coupling of CBF and metabolism). If the MAP falls below 50 mmHg, and in certain pathological states (e.g., ischemia), autoregulation fails and CBF declines. Vascular stenosis or occlusion induces compensatory vasodilatation downstream, which increases the cerebral blood volume and CBF (vascular reserve); the extent of local brain injury depends on the availability of collateral flow, the duration of hemodynamic insufficiency, and the vulnerability of the particular brain region affected. Major neurological deficits arise only when CBF falls below the critical ischemia threshold (ca. 20 ml/100 g/min). Hypoperfusion. If adequate CBF is not reestablished, clinically evident neurological dysfunction ensues (breakdown of cerebral metabolism \Rightarrow EEG and evoked potential change). Prolonged. severe depression of CBF below the infarction threshold of ca. 8-10 ml/100 g/min causes progressive and irreversible abolition of all cellular metabolic processes, accompanied by structural breakdown (necrosis). Infarction occurs where hypoperfusion is most severe; the area of tissue surrounding the zone of infarction in which the CBF lies between the thresholds for ischemia and infarction is called the ischemic penumbra. Brain tissue in the zone of infarction is irretrievably lost, while that in the ischemic penumbra is at risk, but potentially recoverable. The longer the ischemia lasts, the more likely infarction will occur: thus, time is brain. The zones of infarction and ischemia are well demonstrated by recently developed MRI techniques (DWI, PWI).1,2

Stroke Treatment

Primary prevention involves the therapeutic modification or elimination of risk factors.

Special treatment measures: APT (after exclusion of hemorrhage); thrombolysis, treatment of cerebral edema, surgical decompression in space-occupying cerebellar or MCA infarcts, and anticonvulsants, as needed.

Secondary prevention. APT (TIA, mild stroke, atherothrombotic stroke); oral anticoagulation (cardiac embolism, arterial dissection); endarterectomy (in symptomatic carotid stenosis >70%, or >80%, or after mild strokes). The potential utility and indications of carotid angioplasty and stenting in the treatment of carotid stenosis are currently under intensive study.

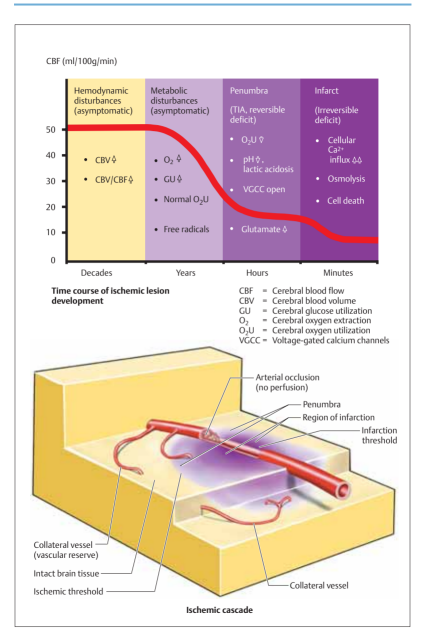
Patients with asymptomatic stenosis are given antiplatelet therapy (APT) consisting of aspirin, aspirin-dipyridamole combination, or clopidogrel. Endarterectomy may be indicated in asymptomatic high-grade stenosis (>80%4 or >90%3). Anticoagulants may be indicated in patients with atrial fibrillation without rheumatic valvular heart disease, depending on their individual risk profile (TIAs, age, comorbidities). Acute treatment is based on the existence of a 3-6-hour interval between the onset of ischemia and the occurrence of maximum irreversible tissue damage (treatment window). General treatment measures include the assurance of adequate cardiorespiratory status (normal blood oxygenation is essential for the survival of the ischemic penumbra); because autoregulation of CBF in the penumbra is impaired, the systolic BP should be maintained above 160 mmHg. The serum glucose level should not be allowed to exceed 200 mg/100 ml. Balanced fluid replacement should be provided, and fever, if it occurs, should be treated. Physicians should be vigilant in the recognition and treatment of complications such as aspiration (secondary to dysphagia), deep venous thrombosis (secondary to immobility of a plegic limb), cardiac arrhythmia, pneumonia, urinary tract infection, and pressure sores. Rehabilitation measures include physical, occupational, and speech therapy, as well as psychological counseling of the patient and family.

¹Diffusion-weighted imaging (DWI) demonstrates the zone of infarction; the early CT signs of infarction (blurring of insular cortex, hypodensity of basal ganglia, cortical swelling) are less reliable.

²Perfusion-weighted imaging (PWI) demonstrates the ischemic penumbra and zone of oligemia (tissue at risk).

³Data from the European Carotid Surgery Trial (ECST).

⁴Data from the North American Symptomatic Carotid Endarterectomy Trial (NASCET).



Clinical Features

Spontaneous (i.e., nontraumatic) intracranial hemorrhage may be epidural, subdural (p. 267), subarachnoid, intraparenchymal, or intraventricular. Its site and extent are readily seen on CT (somewhat less well on MRI) and determine its clinical manifestations.

Subarachnoid Hemorrhage (SAH)

Symptoms and signs. The typical presentation of aneurysmal rupture (by far the most common cause of SAH) is with a very severe headache of abrupt onset ("the worst headache of my life"), often initially accompanied by nausea, vomiting, diaphoresis, and impairment of consciousness. The neck is stiff, and neck flexion is painful. There may also be focal neurological signs, photophobia, and/or backache. Subarachnoid blood can be seen on CT within 24 hours of the hemorrhage in roughly 90% of cases. If SAH is suspected but the CT is negative, a diagnostic lumbar puncture must be performed.

Complications. The initial hemorrhage may extend beyond the subarachnoid space into the brain parenchyma, the subarachnoid space, and/ or the ventricular system. A ruptured saccular aneurysm may rebleed at any time until it is definitively treated; the rebleed risk is highest on the day of onset (day 0), and 40% in the ensuing 4 weeks. The greater the amount of blood in the subarachnoid cisterns, the more likely that vasospasm and delayed cerebral ischemia will occur; the risk is highest between days 4 and 12. Clotted blood blocking the ventricular system or the arachnoid villi can lead to hydrocephalus, of obstructive or malresorptive type, respectively (p. 162). Other complications include cerebral edema, hyponatremia, neurogenic pulmonary edema, seizures, and cardiac arrhythmias.

Intracerebral Hemorrhage

Intraparenchymal hemorrhages of arterial origin are to be distinguished from secondary hemorrhages into arterial or venous infarcts.

General features. Sudden onset of headache, impairment of consciousness, nausea, vomiting, and focal neurological signs, with acute progression over minutes or hours.

Hemorrhage into the basal ganglia. Putaminal hemorrhage produces contralateral hemipare-

sis/hemiplegia and hemisensory deficit, conjugate horizontal gaze deviation, homonymous hemianopsia, and aphasia (dominant side) or hemineglect (nondominant side). *Thalamic hemorrhage* produces similar manifestations and also vertical gaze palsy, miotic, unreactive pupils, and (sometimes) convergence paresis. The very rare *caudate hemorrhages* are characterized by confusion, disorientation, and contralateral hemiparesis. Hemorrhage into the *basal ganglia and internal capsule* leads to coma, contralateral hemiplegia, homonymous hemianopsia, and aphasia (dominant side).

Lobar hemorrhage usually originates at the gray-white matter junction and extends inward into the white matter, producing variable clinical manifestations. Frontal lobe: Frontal head-ache, abulia, contralateral hemiparesis (arm more than leg). Temporal lobe: Pain around the ear, aphasia (dominant side), confusion, upper quadrantanopsia. Parietal lobe: Temporal head-ache, contralateral sensory deficit, aphasia, lower quadrantanopsia. Occipital lobe: Ipsi-lateral periorbital pain, hemianopsia.

Cerebellar hemorrhages are usually restricted to one hemisphere. They produce nausea, vomiting, severe occipital headache, dizziness, and ataxia.

Brain stem hemorrhage. Pontine hemorrhage is the most common type, producing coma, quadriplegia/decerebration, bilateral miosis (pinpoint pupils), "ocular bobbing," and horizontal gaze palsy. Locked-in syndrome may ensue.

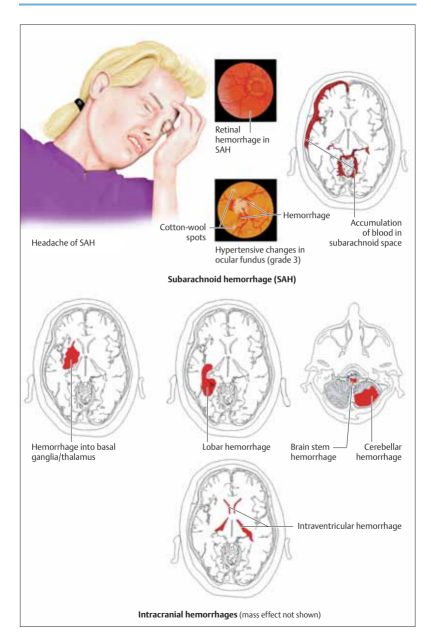
General complications. Intraventricular extension of hemorrhage, hydrocephalus, cerebral edema, intracranial hypertension, seizures, and hemodynamic changes (often a dangerous elevation of blood pressure).

Intraventricular Hemorrhage

Intraventricular hemorrhage only rarely originates in the ventricle itself (choroid plexus). It is much more commonly the intraventricular extension of an aneurysmal SAH or other brain hemorrhage.

Symptoms and signs. Acute onset of headache, nausea, vomiting, impairment of consciousness or coma.

Complications. Extension of hemorrhage, hydrocephalus, seizures.



Pathogenesis

Subarachnoid hemorrhage (SAH). Roughly 85% of cases of SAH are caused by rupture of a saccular aneurysm at the base of the skull. Another 10% are caused by nonaneurysmal lesions (whose nature is not, at present, understood), with bleeding mainly in the perimesencephalic cisterns (p. 8). Other, rare, causes include vertebral artery dissection, arteriovenous malformation (AVM), cavernoma, hypertension, anticoagulation, and trauma.

Aneurysm. An aneurysm is not a congenital lesion, but a progressive, localized dilatation of an arterial wall. Saccular aneurysms tend to develop at branching sites of the internal carotid artery (ICA), the anterior communicating artery, and the proximal middle cerebral artery (MCA). Fusiform aneurysms usually appear as an elongated, twisted, and dilated segment (dolichoectasis) of the basilar artery or supraclinoid ICA. Most are due to atherosclerosis. Spontaneous hemorrhage is rare; ischemia due to arterioarterial embolism is more common. The rare septic-embolic aneurysms (mycotic aneurysms) may be secondary to endocarditis, meningoencephalitis, hemodialysis, or intravenous drug administration. They are located in distal vessel segments, particularly in the MCA, and may cause SAH, massive hemorrhage, or infarction with secondary hemorrhage.

Vascular malformations. Arteriovenous malformations (AVMs) are congenital lesions consisting of a tangled web of arteries and veins with pathological arteriovenous shunting. Most are located near the surface of the cerebral hemispheres. AVMs tend to enlarge over time and may become calcified. They may cause subarachnoid or intracerebral hemorrhage at any age. They become clinically manifest either through a hemorrhage or through headache, seizures, or focal neurological signs (aphasia, hemiparesis, hemianopsia).

Cavernomas are compact, often calcified, aggregations of dilated blood vessels and connective tissue in the brain and leptomeninges. They rarely bleed (ca. 0.5%/year). Some cause seizures and focal deficits; others are discovered on MRI scans as an incidental finding.

Intracerebral hemorrhage. Hypertension is the most common cause; other causes include

aneurysm, AVM, cerebral amyloid angiography, moyamoya, coagulopathy (leukemia, thrombocytopenia, therapeutic anticoagulation), cerebral vasculitis, cerebral venous thrombosis, drug abuse (cocaine, heroin), alcohol, metastases, and brain tumors. Massive hypertensive hemorrhage is thought to be caused by pressure-induced rupture of arterioles and microaneurysms. The high pressure and a kind of chain reaction involving multiple microaneurysms is thought to explain the size of these hemorrhages (despite the small caliber of the ruptured vessels). The recurrent, mainly cortical, bleeding associated with cerebral amyloid angiopathy is due to the fragility of leptomeningeal and cortical small vessels in whose walls amyloid deposits have accumulated.

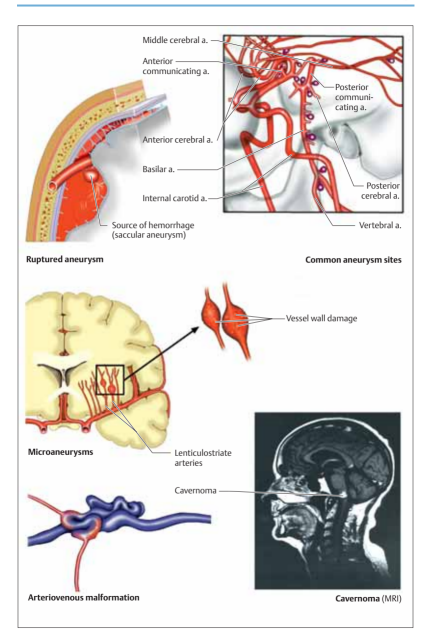
Dural fistula is an abnormal anastomosis between dural arteries and a venous sinus. Hemorrhage is rare; they may cause pulsating tinnitus, headache, papilledema, and visual disturbances.

Treatment

Aneurysmal hemorrhage. Cautious transport, bed rest, analgesia, admission to a neurosurgical unit, and angiography to establish the diagnosis and define the anatomy of the aneurysm(s). The timing of surgical clipping depends on the site and clinical severity of the hemorrhage, the aneurysm's configuration, and the age and general medical condition of the patient. Inoperable cases can be managed with intravascular neuroradiological techniques ("embolization"; filling with Guglielmi detachable coils (GDC) is currently favored).

Hemorrhage from an AVM may be treated with surgery, embolization, and/or stereotactic radiosurgery, depending on its site and extent. Cavernomas that bleed are usually excised.

Intracerebral hemorrhage is often treated conservatively, unless it impairs consciousness or causes a progressive neurological deficit. Major cerebellar hemorrhage (rule of thumb: > 3 cm) is life-threatening unless treated neurosurgically.



Sinus Thrombosis

Symptoms and signs. Aseptic sinus thrombosis most commonly affects the superior sagittal sinus and produces initial headache, vomiting, and focal epileptic seizures, followed by monoparesis or hemiparesis, papilledema, abulia, and impairment of consciousness. Blockage of venous outflow causes cerebral edema and rupture of the distended cerebral veins upstream from the thrombosis. Septic sinus thrombosis is heralded by fever, chills, and malaise. Pain, redness, and swelling of the eye or ear may develop, in addition to focal neurological signs. Transverse and sigmoid sinus thromboses are often secondary to ear and mastoid infections, while cavernous sinus thrombosis is often due to infections about the face (orbit, paranasal sinuses, teeth).

Etiology. Aseptic sinus thrombosis may occur during or after pregnancy, or secondary to the use of oral contraceptives, deficiencies of protein C, protein S, antithrombin III, or factor V, dehydration, polycythemia vera, leukemia, Behçet disease, trauma, surgical procedures, and malignancy. Septic sinus thrombosis often occurs by secondary spread of infections about the head (sinusitis, otitis media, mastoiditis, facial furuncle). *Identification:* CT or MRI angiography generally suffices to demonstrate the occluded sinus; conventional angiography is rarely needed. **Treatment**. *Aseptic thrombosis:* anticoagulation. *Septic thrombosis* (pp. 226, 375): antibiotics and surgical management of the source of infection.

Cerebral Vasculitis

Primary vasculitis arises in the cerebral arteries and veins themselves, while secondary vasculitis is a sequela of another disease (see Causes, below).

Symptoms and signs. Cerebral vasculitis produces variable symptoms and signs, including recurrent ischemia, intracerebral or subarachnoid hemorrhage, persistent headache, focal epilepsy, gradually progressive focal neurological signs, dementia, behavioral abnormalities, cranial nerve palsies, and meningismus. Vasculitis may also affect vessels of the spinal cord (transverse cord syndrome) and those supplying the peripheral nerves (painful mononeuropathy).

Causes. Isolated cerebral angiitis (idiopathic) is difficult to diagnose because its findings are nonspecific (elevated CSF protein, EEG and MRI abnormalities). Leptomeningeal biopsy may be needed. Primary or secondary vasculitides of various kinds affect the vessels of the central nervous system (CNS), peripheral nervous system (PNS) and/or skeletal muscles to a variable extent (see table).

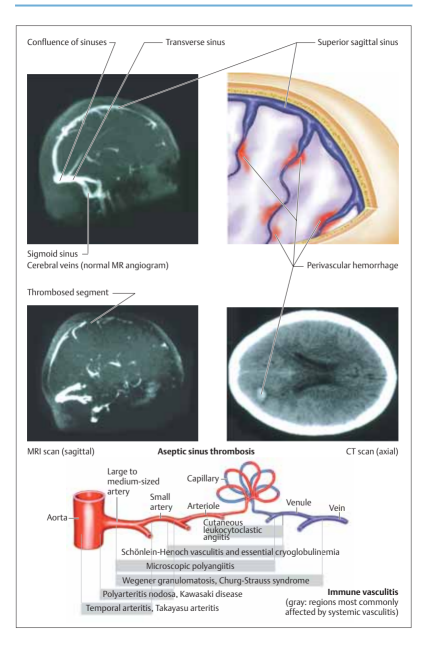
Treatment. Infectious vasculitis is treated with antiviral or antibacterial agents, as needed, while autoimmune vasculitis is treated with corticosteroids and immune suppressants (cyclophosphamide, azathioprine).

Syndrome	CNS	PNS	Muscle
Churg-Strauss syndrome	++	+++	+
Wegener granulomatosis	++	++	+
Behçet disease	++	+/-	-
Lymphomatoid granulomatosis	++	+/-	-
Syphilis, tuberculosis, herpes zoster, bacterial meningitis, fungal infection	++	+/-	+/-
Temporal arteritis	+	+/-	-
Polyarteritis nodosa	+	+++	+
Takayasu arteritis	+	-	-
Lymphoma	+	+	-

180

+++ usual, ++ common, + occasional, +/- rare, - absent

(From Moore and Calabrese, 1994)



Tension Headache

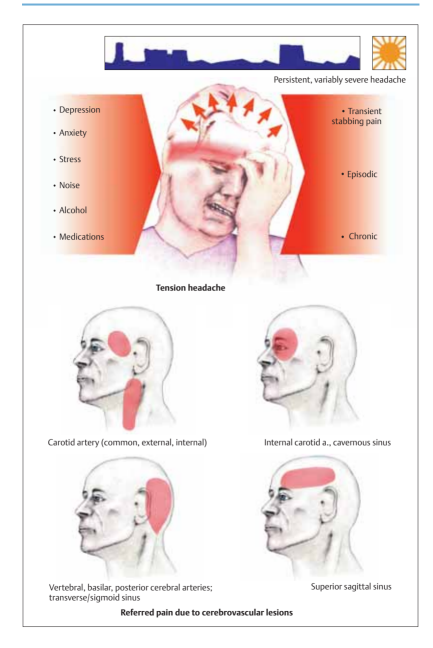
Tension headache often involves painful cervical muscle spasm, may change with the weather, and is frequently ascribed by patients and others to cervical spinal degenerative disease, visual disturbance, or life stress. It consists of a bilateral, prominent nuchal pressure sensation that progresses over the course of the day. Patients may report feeling as if their head were being squeezed in a vise or by a band being drawn ever more tightly around it, or as if their head were about to explode, though the pain is rarely so severe as to impede performance of the usual daily tasks, e.g., at work. It may be accompanied by malaise, anorexia, lack of concentration, emotional lability, chest pain, and mild hypersensitivity to light and noise. Unlike migraine, it is not aggravated by exertion (e.g., climbing stairs), nor does it produce vomiting or focal neurological deficits. It can be episodic (<15 days/month, pain lasting from 30 minutes to 1 week) or chronic (>15 days/month for at least 6 months). Some patients suffer from pericranial tenderness (posterior cervical, masticatory, and cranial muscles). Isolated attacks of sudden, stabbing pain (ice-pick headache) may occur on one side of the head or neck. Tension headache rarely wakes the patient from sleep. Its cause usually cannot be determined, though it may be due to disorders of the temporomandibular joint or psychosomatic troubles such as stress, depression, anxiety, inadequate sleep, or substance abuse. Tension headache combined with migraine is termed combination headache. Pathogenesis. It is theorized that diminished activity of certain neurotransmitters (e.g., endogenous opioids, serotonin) may lead to abnormal nociceptive processing (p. 108) and thus produce a pathological pain state.

Headache Due to Vascular Processes (Other than Migraine)

The nociceptive innervation of the extracranial and intracranial vessels is of such a nature that pain arising from them is often projected to a site in the head that is some distance away from the responsible lesion. Thus, specific diagnostic studies are usually needed to pinpoint the location of the disturbance. The pain may precede the actual vascular event (arterial dissection, arteriovenous malformation, vasculitis), may occur simultaneously with the event (subarachnoid hemorrhage, intracerebral hemorrhage, epidural hematoma, cerebral venous thrombosis, giant cell arteritis, carotidynia, venous outflow obstruction in goiter or mediastinal processes, pheochromocytoma, preeclampsia, malignant hypertension), or may follow the event (subdural hematoma, intracerebral hemorrhage, endarterectomy).

Chronic Daily Headache

Rational treatment is based on the classification of primary daily headache by clinical characteristics (see Table 23, p. 373), and of secondary (symptomatic) headache by etiology.



Migraine

Migraine is a periodic headache often accompanied by nausea and sensitivity to light and noise (photophobia and phonophobia). A typical attack consists of a *prodromal phase* of warning (premonitory) symptoms, followed by an *aura*, the actual *headache phase*, and a *resolution phase*. Attack characteristics often change over time. Attacks often tend to occur in the morning or evening but may occur at any time. They typically last 4–72 hours.

Symptoms and Signs

Prodromal phase. The migraine attack may be preceded by a period of variable prodromal phenomena lasting a few hours to two days. Most patients complain of sensitivity to smells and noise, irritability, restlessness, drowsiness, fatigue, lack of concentration, depression, and polyuria. In children, the chief complaints are abdominal pain and dizziness.

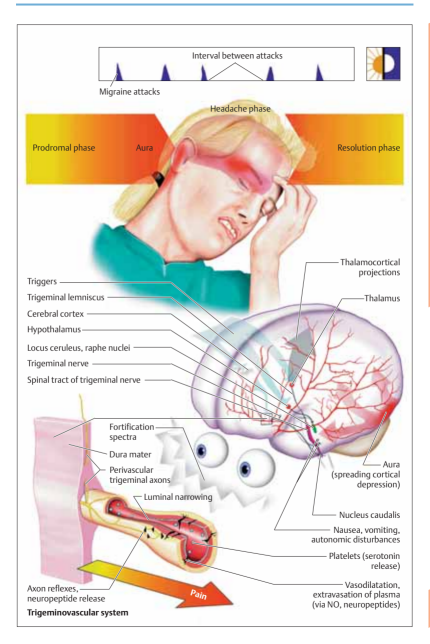
Aura. This is the period preceding the focal cerebral symptoms of the actual migraine headache. Some patients experience attacks without an aura (common migraine), while others have attacks with an aura (classic migraine) that develops over 5-20 minutes and usually lasts less than one hour, but may persist as long as one week (prolonged aura). In some cases, the aura is not followed by a headache ("migraine equivalent"). Auras typically involve visual disturbances, which can range from undulating lines (resembling hot air rising), lightning flashes, circles, sparks or flashing lights (photopsia), or zig-zag lines (fortification figures, teichopsia, scintillating scotoma). The visual images, which may be white or colored, cause gaps in the visual field and usually have scintillating margins. Unilateral paresthesiae (tingling or cold sensations) may occur. Emotional changes (anxiety, restlessness, panic, euphoria, grief, aversion) of variable intensity are relatively common.

Headache phase. Most patients (ca. 60%) complain of pulsating, throbbing, or continuous pain on one side of the head (hemicrania). Others have pain in the entire head, particularly behind the eyes ("as if the eye were being pushed out"), in the nuchal region, or in the temples. Migraine headache worsens on physical exertion and is often accompanied by anorexia, malaise, nausea, and vomiting.

Resolution phase. This phase is characterized by listlessness, lack of concentration, and increased pain sensitivity in the head.

Pathogenesis

During the interval between attacks, various disturbances (genetically determined) may be observed, e.g., cerebral hypomagnesemia, elevated concentration of excitatory amino acids (glutamate, aspartate), and increased reactivity of cranial blood vessels. The cumulative effect of these disturbances is a heightened sensitivity to nociceptive stimuli (migraine pain threshold). Impulses from the cortex, thalamus, and hypothalamus activate the so-called migraine center responsible for the generation of migraine attacks, putatively located in the brain stem (serotonergic raphe nuclei, locus ceruleus). The migraine center triggers *cortical spreading* depression (suppression of brain activity across the cortex) accompanied by oligemia, resulting in an aura. Trigeminovascular input from meningeal vessels is relayed to the brain stem, via projecting fibers to the thalamus and then, by the parasympathetic efferent pathway, back to the meningeal vessels (trigeminal autonomic reflex circuit). Perivascular trigeminal C-fiber endings (trigeminovascular system) are stimulated to release vasoactive neuropeptides such as substrate P, neurokinin A, and calcitonin gene-regulated polypeptide (CGRP), causing a (sterile) neurogenic inflammatory response. Vasoconstriction and vascular hyperesthesia with subsequent vasodilatation spread via trigeminal axon reflexes. The perception of pain is mediated by the pathway from the trigeminal nerve to the nucleus caudalis, thalamus (p.94) and cortex. Trigeminal impulses also reach autonomic centers.



Trigeminal Neuralgia

Trigeminal neuralgia (tic douloureux) is characterized by the sudden onset of excruciating, intense stabbing pain (during waking hours). Several brief attacks (<2 minutes each) generally occur in succession. The pain is almost always precipitated by a trigger stimulus or activity (e.g., chewing, speaking, swallowing, touching the face, cold air, tooth brushing, shaving) and is located in the distribution of one or two branches of the trigeminal nerve, usually V/2 and/or V/3. Involvement of V/1, all three branches, or both sides of the face is uncommon. The attacks may persist for weeks to months or may spontaneously remit for weeks, or even vears, before another attack occurs, Trigeminal neuralgia in the V/3 distribution is often mistaken for odontogenic pain, sometimes resulting in unnecessary tooth extraction. Typical (idiopathic) trigeminal neuralgia must be distinguished from secondary forms of the syndrome (see below).

Pathogenesis. Idiopathic trigeminal neuralgia \Rightarrow much evidence points to microvascular compression of the trigeminal nerve root (usually by a branch of the superior cerebellar artery) where it enters the brain stem, leading to the development of ephapses or suppression of central inhibitory mechanisms. *Symptomatic trigeminal neuralgia* \Rightarrow cerebellopontine angle tumors, multiple sclerosis, vascular malformations.

Cluster Headache (CH)

Episodic cluster headache. Attacks of very severe burning, searing, stabbing, burning, needlelike, or throbbing pain develop over a few minutes on one side of the head, behind or around the eye, and may extend to the forehead, temple, ear, mouth, jaw, throat, or nuchal region. If untreated, attacks last ca. 15 minutes to 3 hours. They are predominantly nocturnal, waking the patient from sleep, but can also occur during the day. Attacks come in episodes (clusters) consisting of 1–3 daily bouts of pain for up to 8 weeks. A seasonal pattern of occurrence may be observed. During a cluster, the pain can be triggered by alcoholic drinks, histamines, or nitrates. Temporal pressure or the

application of heat to the eye may alleviate the pain. Unlike migraine patients, who seek peace and quiet, these patients characteristically pace restlessly, and may even strike their aching head with a fist. The headache may be accompanied by ipsilateral *ocular* (watery eyes, conjunctival injection, incomplete Horner syndrome, photophobia), *nasal* (nasal congestion, rhinorrhea), and *autonomic* manifestations (facial flushing, tenderness of temporal artery, nausea, diarrhea, polyuria, fluctuating blood pressure, cardiac arrhythmia). Cluster headache is more common in men.

Chronic cluster headache. Attacks do not occur in clusters, but rather persist for more than one year at a time, punctuated by remissions lasting no longer than two weeks. Chronic cluster headache may arise primarily, or else as a confluence of clusters in what began as episodic cluster headache.

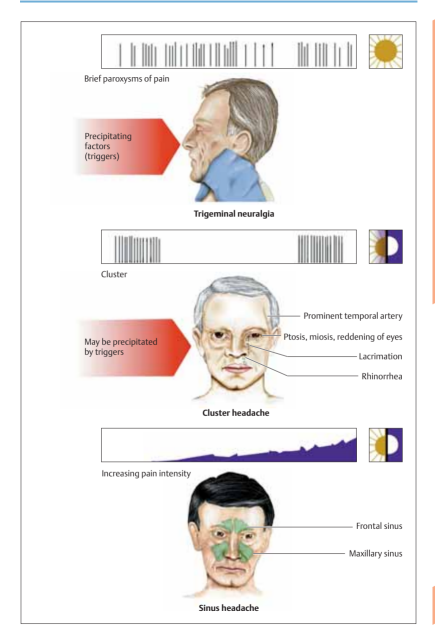
Pathogenesis. One hypothesis attributes CH to dilatation of the carotid artery within the carotid canal, causing compression of the periarterial sympathetic plexus. There is also evidence suggesting a role for inflammatory dilatation of the intracavernous venous plexus. The result is abnormal function of the sympathetic and parasympathetic fibers in the region of the cavernous sinus (□ autonomic dysfunction, activation of trigeminovascular system).

Chronic Paroxysmal Hemicrania (CPH)

CPH is a very rare condition characterized by the daily onset of pain similar to that of cluster headache. The daily attacks of CPH are much more frequent (10–20 times/day) and shorter (5–30 minutes) than those of cluster headache. The pain of CPH typically responds to indomethacin.

Sinus Headache

The pain of frontal, sphenoid, or ethmoid nasal sinusitis is usually felt in the middle of the forehead and above the eyes. That of maxillary sinusitis radiates to the upper jaw and zygomatic region and worsens when the patient bends forward.



Nociceptive Transmission

The brain tissue itself is insensitive to pain. The major cranial and proximal intracranial vessels and dura mater of the supratentorial compartment derive nociceptive innervation from the ophthalmic nerve (V/1, p. 94), while those of the posterior fossa are innervated by C2 branches. Because nociceptive impulses from the anterior and middle fossae, the venous sinuses, the falx cerebri, and the upper surface of the tentorium travel through V/1, the pain that is experienced is referred to the ocular and frontoparietal regions; similarly, pain arising from the lower surface of the tentorium, the posterior cranial fossa, and the upper 2-3 cervical vertebrae (mediated by C2) is referred to the occipital and nuchal region. Small regions of the dura mater are innervated by CN IX and X; pain arising here is, accordingly, referred to the throat or ear. These neuroanatomical connections also explain the referral of pain from the upper cervical region to the eye (shared trigeminal innervation), and why tension and migraine headache can cause pain in the neck.

Cervical Syndrome (Upper Cervical Syndrome)

Cervical syndrome typically causes pain in the frontal, ocular, and nuchal regions. The pain is usually continuous, without any circadian pattern, but may be more severe during the day or night. It may be worsened by active or passive movement of the head. It is usually due to a lesion affecting the C2 root and is characterized by muscle spasm, tenderness, and restricted neck movement. The diagnosis is based on the typical clinical findings, and cannot be based solely on radiographic evidence of degenerative disease of the cervical spine. For other causes of neck pain, see Table 23 (p. 373). For cervical distortion (whiplash), see p. 272. For *Posttraumatic headache*, see p. 270.

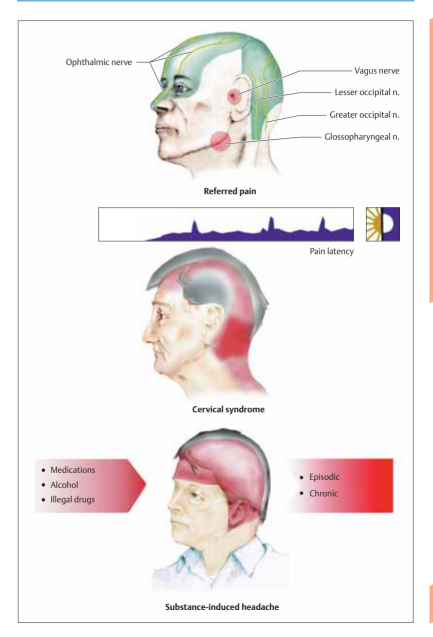
Substance-Induced Headache

Acute headache can be induced by a number of vasoactive substances. Triggers include alcohol consumption or withdrawal ("hangover"), caffeine or nicotine withdrawal, sodium glutamate,

cocaine, marijuana, nitrates, and dihydropyridines (calcium antagonists). The headache is usually a pressing, piercing, or pulsating pain, and is typically bifrontal or frontotemporal. It may be accompanied by nausea, chest tightness, dizziness, abdominal complaints, lack of concentration, or impairment of consciousness.

Rebound headache. Persons suffering from recurrent or chronic headache are at risk for the excessive or uncontrolled use of medications. singly or in combination (analgesics, benzodiazepines, ergot alkaloids, combined preparations). This may result in daily rebound headache, persisting from morning to night and characterized by pressurelike or pulsating, unilateral or bilateral pain, accompanied by malaise, nausea, vomiting, phonophobia, and photophobia. Patients may also complain of lack of concentration, disturbed sleep, blurred or flickering vision, a feeling of cold, and mood swings. These patients change medications frequently and tend to take medication even at the first sign of mild pain, because they fear a recurrence of severe pain. Eventually, drug tolerance develops, resulting in persistent headache. The original migraine or tension headache may be largely masked by the rebound headache. Other drug side effects may include ergotism, gastritis, gastrointestinal ulcers, renal failure, physical dependence, and epileptic seizures (withdrawal seizures).

Central Nervous System



Treatment

The proper treatment of headache depends on its cause. Episodic or chronic tension headache and migraine are by far the most common types of headache. Structural lesions are a rare cause of headache (< 5% of all headaches); such headaches typically start suddenly, worsen quickly, and represent a new type of pain that the patient has never had before. If a structural lesion is suspected, neuroimaging studies should be performed.

General Treatment Measures

Good patient-physician communication is essential for the diagnosis and treatment of headache. The most important clues to differential diagnosis are derived from the case history. The medication history must be obtained, and psychological factors must also be considered; headache is often associated with anxiety, e.g., fear of a brain tumor, of a mental illness, or of not having one's complaints taken seriously. Patients must be instructed how they themselves can improve their symptoms (behavioral modification) through lifestyle changes (e.g., avoidance of alcohol, dietary changes, physical exercise, adequate sleep) and nonpharmacological measures (relaxation training, biofeedback, stress management, keeping a pain diary).

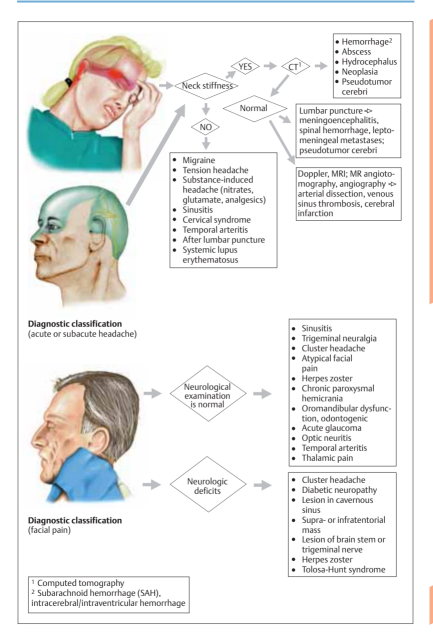
Acute Treatment

Type of Headache	General Measures	Pharmacotherapy
Episodic tension headache	Behavioral therapy, ice packs	Peppermint oil to forehead and temples; aspirin, acetaminophen, ibuprofen, or naproxen
Migraine	Rest, ice packs	Antiemetic (metoclopramide or domperi- done) + aspirin or acetaminophen; if ineffec- tive, triptans ¹
Cluster headache	Hot compresses	Oxygen inhalation; if ineffective, triptan s.c. or ergotamine
Chronic paroxysmal hemicrania		Indomethacin
Trigeminal neuralgia	Avoidance of triggers	Carbamazepine, gabapentin, phenytoin, ba- clofen, or pimozide ²

This group includes sumatriptan, almotriptan, eletriptan, frovatriptan, naratriptan, rizatriptan, and zolmitriptan.
 Patients with primary or secondary resistance to medical treatment should be treated neurosurgically (percutaneous thermocoagulation or retroganglionic glycerol instillation; microvascular decompression).

Prophylaxis

Type of Headache	General Measures	Pharmacotherapy
Episodic or chronic tension headache	Behavioral therapy	Tricyclic antidepressants (e.g., amitryptiline, doxepin, amitryptiline oxide)
Migraine	Behavioral therapy	1st line: Beta-blockers (e.g., metoprolol, pro- pranolol); 2nd line: flunarizine or valproate; 3rd line: methysergide or pizotifen
Episodic cluster headache	Avoidance of alcohol (during cluster), nitrates, histamines, and nicotine	Prednisone, ergotamine, verapamil, methy- sergide, or lithium
Chronic cluster headache		Lithium, verapamil, or pizotifen



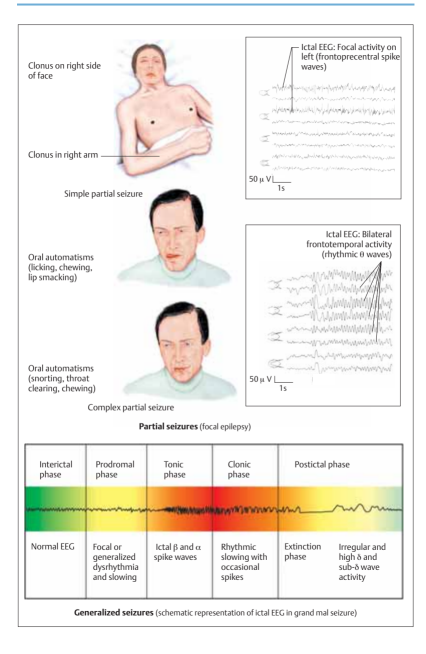
An epileptic seizure (convulsion, fit) is a sign of brain dysfunction (p. 198). Seizures generally last no more than 2 minutes; the *postictal period* may be marked by impairment of consciousness or focal neurological signs. The type and extent of motor, sensory, autonomic and/or psychological disturbance during the seizure (seizure semiology) reflects the location and extent (localized/generalized) of brain dysfunction. Seizure classification, including the differentiation of true epileptic from nonepileptic seizures (pseudoseizures or psychogenic seizures, p. 202), is essential for effective treatment.

Partial (Focal) Seizures

Focal or partial seizures reflect paroxysmal discharges restricted to a part of the affected hemisphere. By definition, *simple partial seizures* are those in which consciousness is not impaired, while *complex partial seizures* (psychomotor seizures) are those in which consciousness is impaired. A sensory or behavioral disturbance preceding a focal or generalized seizure with motor manifestations is called a seizure *aura*. Some features of partial seizures are listed in the table below. The semiology of simple and complex partial seizures depends on their site of origin (focus) and the brain areas to which they spread (see table, p. 194). They may become *secondarily generalized*, evolving into generalized paroxysmal attacks or tonic-clonic seizures. The initial symptoms and signs vary depending on the location of the epileptic focus.

Feature	Simple Partial Seizures	Complex Partial Seizures
Consciousness	Unaffected	Impaired
Duration	Seconds to minutes	Minutes
Symptoms and signs	Depend on site of origin; no postictal confusion	Depend on site of origin; postictal confusion
Age group	Any age	Any age
Ictal EEG	Contralateral epileptiform discharges; in many cases, no interictal abnormali- ties are detected	Unilateral or bilateral epileptiform discharges, diffuse or focal

(Adapted from Gram, 1990)



Site of focus	Seizure type	Symptoms and signs
Frontal lobe	Simple or complex par- tial seizures with or without secondary generalization (hypermo- tor frontal lobe seizures)	Adversive head movement and other complex motor phenomena (mainly in legs), e. g., pelvic movements, ambulatory automatisms, swimming movements, fencing posture, staring, laughing, outcries, genital fumbling, autonomic dysfunction, speech arrest. Mood changes may also occur. Seizures occur several times a day with an abrupt onset. The unvarying course of the seizures may suggest hysteria. Brief postictal confusion
Temporal lobe	Complex partial seizures with or without second- ary generalization	Ascending epigastric sensations (nausea, heat sensa- tion); olfactory/gustatory hallucinations; compulsive thoughts, feeling of detachment, déjà-vu, jamais-vu; oral and other automatisms (psychomotor attacks); dyspnea, urinary urgency, palpitations; macropsia, micropsia; postictal confusion
Parietal lobe	Simple partial seizures with or without second- ary generalization	Sensory and/or motor phenomena (jacksonian seizures); pain (rare)
Occipital lobe	Simple partial seizure with or without second- ary generalization	Unformed visual hallucinations (sparks, flashes)

(Adapted from Gram, 1990)

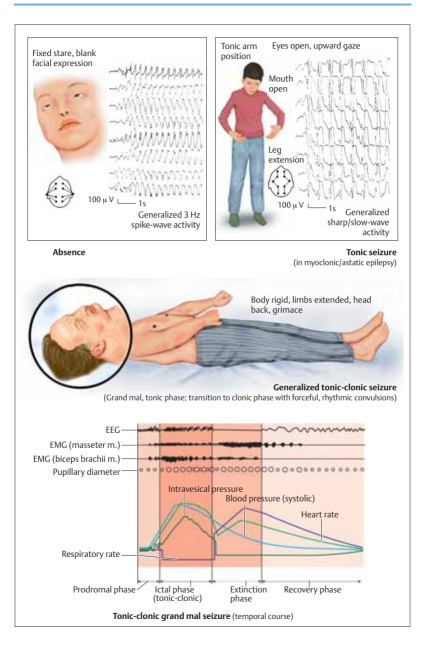
Generalized Seizures

Generalized epilepsy reflects paroxysmal discharges occurring in both hemispheres. The seizures may be either *convulsive* (e.g., generalized tonic-clonic seizure \Rightarrow GTCS) or *nonconvulsive* (myoclonic, tonic, or atonic seizures; absence seizures). Generalized seizures are classified by their clinical features.

Feature	Absence Seizure	Myoclonic Seizure	Atonic (Astatic) Seizure	Tonic-clonic Seizure
Conscious- ness	Impaired	Unaffected	Impaired	Impaired
Duration	A few (\leq 30) seconds	1–5 seconds	A few sec- onds	1–3 minutes
Symptoms and signs	Brief absence, vacant gaze and blinking fol- lowed by immediate return of mental clar- ity; automatisms (lip smacking, chewing, fiddling, fumbling) may occur	Sudden, bi- laterally syn- chronous jerks in arms and legs; often occur in series	Sudden loss of muscle tone causing severe falls	Initial cry (occasionally); falls (loss of muscle tone); respira- tory arrest; cyanosis; tonic, then clonic seizures; muscle re- laxation followed by deep sleep. Tongue biting, urinary and fecal incontinence
Age group	Children and adoles- cents	Children and adolescents	Infants and children	Any age
Ictal EEG	Bilateral regular 3 (2–4) Hz spike waves	Polyspike waves, spike waves, or sharp and slow waves	Polyspike waves, flat- tening or low- voltage fast activity	Often obscured by muscle arti- facts

194

(Adapted from Gram, 1990)



The etiology and prognosis of epilepsy depend on its clinical type. All forms of epilepsy (e.g., absence epilepsy of childhood, juvenile myoclonic epilepsy, temporal lobe epilepsy, frontal lobe epilepsy, reflex epilepsy) are characterized by recurrent paroxysmal attacks; thus classification cannot be based on a single seizure. Epileptic syndromes vary in seizure pattern, cause, age at onset, precipitating factors, EEG changes, and prognosis (e.g., neonatal convulsions, infantile spasms and salaam seizures = West syndrome, Lennox-Gastaut syndrome, temporal lobe seizures). Seizures triggered by fever, substance abuse, alcohol, eclampsia, trauma, tumor, sleep deprivation, or medications are designated as isolated nonrecurring seizures or acute epileptic

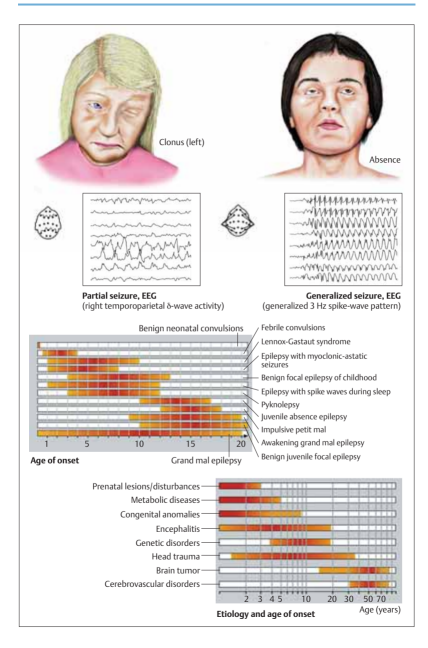
reactions. Status epilepticus is a single prolonged seizure or a series of seizures without full recovery in between. Any type of seizure (convulsive or nonconvulsive) may appear under the guise of status epilepticus. In grand mal status epilepticus, patients do not regain consciousness between seizures.

Location-related (focal, partial) epilepsy can be differentiated from generalized epilepsies and epileptic syndromes on the basis of the seizure pattern. Seizures that cannot be classified because of inadequate data on focal or generalized seizure development are called unclassified epilepsy or epileptic syndrome. Other terms used in classification refer to seizure etiology (e.g., idiopathic, cryptogenic, symptomatic).

Type of Epilepsy	Features
Location-related	Partial (focal) seizures
Generalized	Generalized convulsive seizures (GCS)
Idiopathic	No known cause other than genetic predisposition. No manifestations other than epileptic seizures. Characteristic age of onset
Cryptogenic	Assumed to reflect a CNS disorder of unidentified type. (Once the cause is identified, epilepsy is classified as symptomatic.)
Symptomatic	Due to an identified CNS disorder or lesion

Type of Epilepsy	Etiology	Epilepsy/Epileptic Syndrome
Location-related (focal, localized, partial)	Idiopathic (charac- teristic age of onset)	Benign epilepsy of childhood with centrotemporal spikes; epilepsy of childhood with occipital paroxysms
	Cryptogenic or symptomatic	Variable expression depending on cause and location (e.g., tem- poral, frontal, parietal, or occipital lobe epilepsy)
Generalized	Idiopathic (charac- teristic age of onset)	Absence epilepsy of childhood (pyknolepsy); juvenile absence epilepsy; juvenile myoclonic epilepsy (impulsive petit mal); awak- ening grand mal epilepsy (GTCS); epilepsy with specific triggers (reflex epilepsy)
		West syndrome (infantile spasms, salaam seizures); Lennox– Gastaut syndrome; myoclonic-astatic epilepsy; epilepsy with myoclonic absence
	Symptomatic	Early myoclonic encephalopathy (unspecific etiology); seizures secondary to various diseases
Unsure whether focal or general- ized	Idiopathic or symp- tomatic	Neonatal convulsions; acquired epileptic aphasia (Landau– Kleffner syndrome)
Variably focal and generalized	Symptomatic (situation-related seizure)	Febrile convulsions; isolated seizure or isolated status epilepti- cus; acute metabolic or toxic triggers

Epilepsy: Classification



Causes. Some patients have a *genetic* predisposition to epilepsy, particularly those with generalized epilepsies. Some hereditary diseases are associated with epilepsy (e.g., tuberous sclerosis, Sturge–Weber syndrome, mitochondrial encephalopathies, sphingolipidoses). Acquired forms of epilepsy may be focal (possibly with secondary generalization), bilateral, or diffuse (primary generalized epilepsies). The causes include developmental disorders, pyridoxine deficiency, hippocampal sclerosis, brain tumors, head trauma, cerebrovascular disturbances, alcohol, drug abuse, medications, and CNS infections.

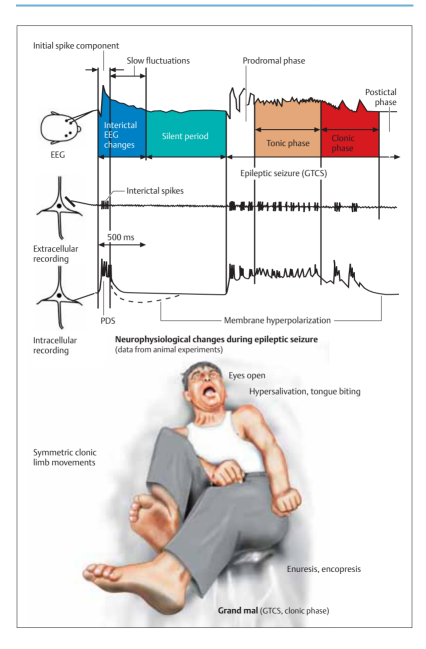
Pathophysiology. Seizure activity in the brain is thought to be initiated by a preponderance of excitatory over inhibitory postsynaptic potentials (EPSP, IPSP), resulting in depolarization of nerve cell membranes. Such a depolarization may appear on the EEG as an interictal spike, an initial spike component, or an abrupt depolarization with superimposed high-frequency action potentials (paroxysmal depolarization shift, PDS). The synchronous discharges of large numbers of neurons result in an epileptic seizure. Seizure activity is terminated by active processes such as transmembrane ion transport via sodium-potassium pumps, adenosine release, and the liberation of endogenous opiates, whose combined effect is membrane hyperpolarization, manifested as slow-wave activity in the EEG. Factors favoring the development of seizures include changes in the concentration of electrolytes (Na⁺, K⁺, Ca²⁺), excitatory amino acids (glutamic acid), and inhibitory amino acids (GABA), irregular interneuron connections, and abnormal afferent connections from subcortical structures (⇒ diencephalon, thalamus, brain stem). In focal epilepsy, the epileptic focus is surrounded by an "inhibitory margin", while the paroxysmal activity of generalized epilepsy is spread throughout the brain. General treatment measures. Lifestyle changes (sleep-wake rhythm, avoidance of seizure triggers); chronic use of anticonvulsant medication. Patients with partial seizures preceded by long auras may be able to abort their seizures while still in the aura phase by various concentration techniques (seizure interruption methods).

Antiepileptic drugs (AEDs). AEDs work by a variety of mechanisms, e.g., inhibition of vol-

tage-gated sodium channels (carbamazepine, oxcarbazepine, lamotrigine, phenytoin, valproic acid) or thalamic calcium channels (ethosuximide), or interaction with inhibitory GABA re-(benzodiazepines, ceptors phenobarbital. gabapentin, tiagabine, levetiracetam) or excitatory glutamate receptors (phenobarbital, felbamate, topiramate). Antiepileptic therapy is generally started in patients who have had a single seizure and are thought to be at risk of recurrence, in those with an epileptic syndrome, and in those who have had two or more seizures within 6 months. AEDs used to treat focal, unclassified. and symptomatic tonic-clonic seizures include carbamazepine, gabapentin, lamotrigine, oxcarbazepine, topiramate, levetiracetam, phenytoin, phenobarbital, and primidone; those used to treat generalized seizures include valproic acid, ethosuximide (absences), primidone, phenobarbital (epilepsy associated with myoclonic seizures, tonic-clonic seizures), and lamotrigine. Treatment is always begun with a single drug (monotherapy); if this ineffective, another drug is used instead of or in addition to the first (combination therapy). Antiepileptic therapy can be discontinued in some cases if the risk of seizure recurrence is judged to be low.

Other measures. Surgery (indicated in patients with drug-resistant focal epilepsy and/or resectable lesions, such as brain tumors or unilateral mesial temporal sclerosis). Vagus nerve stimulation by means of an implanted neurocybernetic prosthesis (NCP) is a form of treatment whose efficacy remains controversial.

Prognosis (Table 24, p. 373). Antiepileptic drugs prevent seizure recurrence in roughly 70% of patients, reduce the frequency of seizures in 25%, and are ineffective in 5% (drug resistance), especially those with Lennox–Gastaut syndrome, symptomatic myoclonic epilepsy, and cryptogenic syndromes.



Nonepileptic Seizures

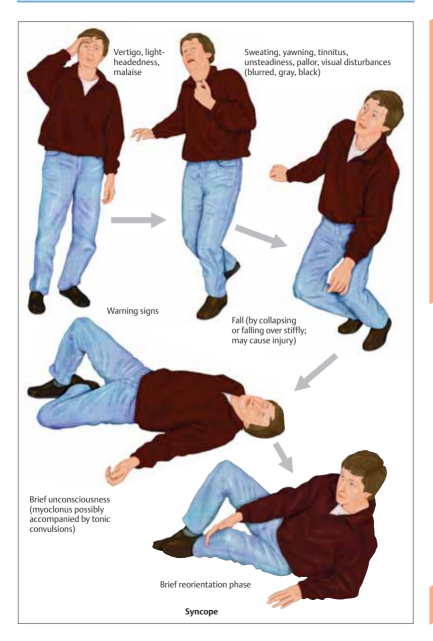
The differentiation of epileptic from nonepileptic seizures is of major prognostic and therapeutic importance. Nonepileptic seizures may or may not involve loss of consciousness. *Pseudoseizures* resemble epileptic seizures (p. 192 ff), but are of nonepileptic origin. This broad category includes syncope, psychogenic seizures, and simulated seizures. "Pseudoseizure," in the narrower sense, is a synonym for "psychogenic seizure."

Nonepileptic seizures are misdiagnosed as epileptic seizures in nearly 20% of patients, roughly 15% of whom are unnecessarily treated with antiepileptic drugs; conversely, some 10% of all epileptic seizures are misdiagnosed as nonepileptic seizures; 20–30% of patients have both epileptic and nonepileptic seizures. In case of doubt, the patient should be referred to a specialist or specialized epilepsy center.

Syncope

Syncope is defined as a brief loss of consciousness, often involving a fall, due to transient cerebral ischemia or hypoxia (see Table 25, p. 374 for potential causes). In 45% of cases, the cause can be determined from the history and physical examination. Important anamnestic clues include triggers such as excitement or anxiety, precipitating situations (blood drawing, prolonged standing, urination, coughing fits, pain), heart disease, mental illness (generalized anxiety disorder, depression, somatization disorders), and medications. The patient should be evaluated for possible blood pressure abnormalities and for cardiac or neurological disorders (p. 148). EEG yields the diagnosis in only about 2% of cases. Only rare cases of syncope are due to TIA (p. 166). Syncope clinically resembles an epileptic seizure in some ways, but differs in others (see table, below).

Clinical Feature	Syncope	Epileptic Seizure
Triggers	Common	No
Time of day	Mostly diurnal; does not awaken patient from sleep	Day or night; awakens patient from sleep
Skin coloration	Pale	Cyanotic or normal
Premonitory symptoms	Tinnitus, visual blurring or blackout, feeling faint, lightheadedness	None or aura
Type of fall	Collapse or fall over stiffly (often back- wards)	Fall over stiffly
Duration	Usually < 30 seconds	1–3 minutes or longer
Abnormal movements (myoclonus)	Frequent, arrhythmic, multifocal to generalized, last < 30 seconds	Always generalized, 1–2 minutes
Eyes	Open	Closed
Urinary incontinence	Occasional	Common
Postictal confusion	Brief or absent	Longer-lasting
Tongue-biting	Occasional	Common
Prolactin, creatine kinase	Normal	Elevated
Typical EEG changes	Absent	Common
Focal neurological deficit	Absent	Occasional



Psychogenic Seizures

Nonorganic, nonepileptic seizures arising from psychological factors do not involve loss of consciousness. They are involuntary and unintentional, and thus must be differentiated from simulated seizures, which are voluntarily, consciously, and intentionally produced events. Psychogenic seizures may resemble frontal lobe seizures (p. 194) and are more common in women than men. About 40% of patients with psychogenic seizures also suffer from true epileptic seizures. The case history often reveals characteristic risk factors, which may be biographical (family difficulties, abuse, divorce, sexual assault in childhood), somatic (genetic predisposition), psychiatric (conflicts, stress, psychosocial gain from illness behavior, mental illness), or social (poor living and working conditions). Patients often meet the psychiatric diagnostic criteria for a conversion disorder (F44.5 according to the ICD-10). Epileptic seizures in family members, or in the patients themselves, may serve as the prototype for psychogenic seizures.

Premonitory signs. Psychogenic seizures can be induced or terminated by suggestion. They may be preceded by a restless, anxious, or fearful state. They usually occur in the presence of others (an "audience") and do not occur when the patient is asleep.

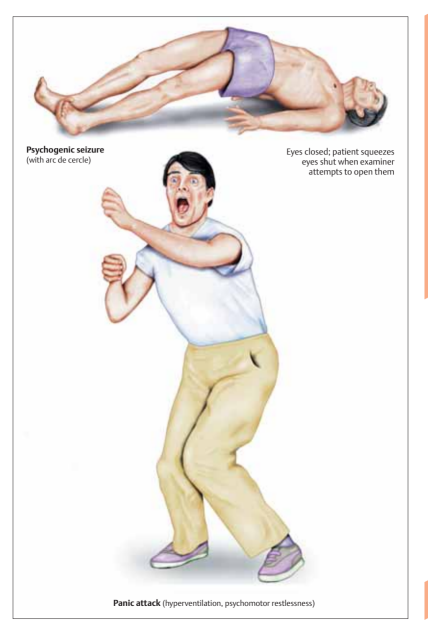
Seizure semiology. Psychogenic seizures usually take a dramatic course, with a variable ending. Their semiology is usually of a type more likely to incite sympathy and pity in onlookers than fear or revulsion. Typical features include an abrupt fall or slow collapse, jerking of the limbs, tonic contraction of the body, writhing (arc de cercle), calling out, shouting, rapid twisting of the head and body, and forward pelvic thrusting; the sequence of movements is usually variable. The eyes are usually closed, but sometimes wide open and staring; the patient squeezes the eyes shut when passive opening is attempted. Urinary incontinence or injury (self-mutilation) may also occur. Tongue-bite injuries, if present, are usually at the tip of the tongue (those in true epileptic fits are usually lateral). The patient is less responsive than normal to external stimuli. including painful stimuli, but not unconscious (squeezes eyelids shut when the eyes are

touched, drops arm to the side when it is held over the patient's face and released). The patient's skin is not pale or cyanotic during the ictus. Patients who hyperventilate during psychogenic seizures may have carpopedal spasms. Psychogenic seizures often last longer than epileptic seizures.

Postictal phase. No focal neurological deficits can be detected, though there may be a psychogenic postictal stupor. The serum prolactin level is not elevated (which, however, does not rule out a true epileptic seizure). The seizure may be terminated abruptly by suggestion, or by departure of the "audience." Some patients recall the seizure to some extent, while others emphatically depy memory of it.

■ Panic Disorder (ICD-10 ⇒ F41.0)

Panic disorder is characterized by sudden, unexpected and apparently unprovoked attacks of intense anxiety, which may range in severity from a general feeling of restlessness to a mortal dread. The attacks usually last 5-30 minutes and may awaken the patient from sleep. Accompanying symptoms include feelings of detachment from the environment, i.e., depersonalization (detachment from one's own body, floating state) and derealization (sensation of being in a dream or nightmare, feeling of unreality); autonomic and other physical symptoms of variable severity, including cardiovascular (tachycardia, palpitations, pallor, chest pain or pressure), gastrointestinal (nausea, dry mouth, dysphagia, diarrhea), respiratory (hyperventilation, dyspnea, smothering sensation), and other manifestations (tremor, twitching of the limbs, dizziness, paresthesia, mydriasis, urinary urgency, sweating). The differential diagnosis includes epilepsy (aura, simple partial seizures), hyperthyroidism. hyperventilation syndrome, pheochromocytoma, heart disease, and hypoglycemia.



Drop Attacks

Sudden, unprovoked, and unheralded falls without loss of consciousness are most common in patients over 65 years of age. Some 10–15% of these drop attacks cause serious injury, particularly fractures. The patient may not be able to get up after a fall. The common causes of recurrent falls in each age group are listed in Table 26 (p. 374). Those associated with loss of consciousness are described on pages 192 ff and 200.

Hyperventilation Syndrome (Tetany)

The clinical manifestations include paresthesiae (perioral, distal symmetrical or unilateral), generalized weakness, palpitations, tachycardia, dry mouth, dysphagia, dyspnea, yawning, pressure sensation in the chest, visual disturbances, tinnitus, dizziness, unsteady gait, muscle stiffness, and carpopedal spasms. The patients report feelings of restlessness, panic, unreality, or confusion. Psychological causes include anxiety. hysteria, and inner conflict. Metabolic causes include hypocalcemia (due to hypoparathyroidism, vitamin D deficiency, malabsorption, or pancreatitis) and a wide range of other disturbances including hypercalcemia, hypomagnesemia, prolonged vomiting, pulmonary embolism, salicylate intoxication, acute myocardial infarction, severe pain, high fever due to septicemia, pneumothorax, stroke, and neurogenic pulmonary edema. Chronic hyperventilation syndromes are more common than acute syndromes, but also more difficult to diagnose.

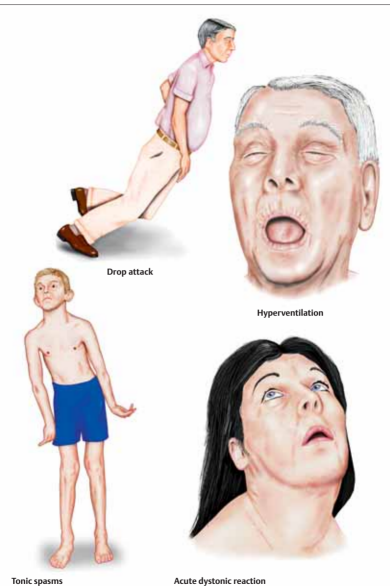
Tonic Spasms

These are unilateral muscular spasms (often painful) that are not accompanied by loss of consciousness; they last seconds to minutes, and occur up to 30 or more times a day. They are most commonly seen in multiple sclerosis, less commonly in cerebrovascular disorders. These spasms are often triggered by movement. Some patients have paresthesiae (tingling, burning) contralateral to the affected side before the muscle spasm sets in. The underlying lesion may be in the brain stem (pons) or internal capsule.

Acute Dystonic Reaction

Acute dystonic reactions can occur within a few hours to one week of starting treatment with dopamine receptor antagonists, e.g., *neuroleptics* (benperidol, fluphenazine, haloperidol, triflupromazine, perphenazine), *antiemetics* (metoclopramide, bromopride), and *calcium antagonists* (flunarizine, cinnarizine). *Symptoms and signs:* focal or segmental dystonia (p. 64), sometimes painful, marked by oculogyric crisis, blepharospasm, pharyngospasm with glossospasm and laryngospasm, or oromandibular dystonia with tonic jaw and tongue movements. Generalized reactions are also seen on occasion (p. 66).

Nonepileptic Seizures



(muscular spasms on left)

Acute dystonic reaction (oculogyric crisis, oromandibular/pharyngeal dystonia) The diagnosis of Parkinson disease (PD; sometimes termed idiopathic Parkinson disease, to distinguish it from symptomatic forms of parkinsonism, and from other primary forms) is mainly based on the typical neurological findings, their evolution over the course of the disease, and their responsiveness to levodopa (Ldopa). Longitudinal observation may be necessary before a definitive diagnosis of PD can be given. PD is characterized by a number of disturbances of motor function (*cardinal manifestations*) and by other *accompanying manifestations* of different kinds and variable severity.

Cardinal Manifestations

Bradykinesia, hypokinesia, and akinesia. Motor disturbances include slow initiation of movement (akinesia), sluggishness of movement (bradykinesia) and diminished spontaneous movement (hypokinesia); these terms are often used nearly interchangeably, as these disturbances all tend to occur together. Spontaneous fluctuations of mobility are not uncommon. The motor disturbances are often more pronounced on one side of the body, especially in the early stages of disease. They affect the craniofacial musculature to produce a masklike facies (hypomimia), defective mouth closure, reduced blinking, dysphagia, salivation (drooling), and speech that is diminished in volume (hypophonia), hoarse, poorly enunciated, and monotonous in pitch (dysarthrophonia). The patient may find it hard to initiate speech, or may repeat syllables; there may be an involuntary acceleration of speech toward the end of a sentence (festination). Postural changes include stooped posture, a mildly flexed and adducted posture of the arms, and postural instability. Gait disturbances appear in the early stages of disease and typically consist of a small-stepped gait, shuffling, and limping, with reduced arm swing. Difficulty initiating gait comes about in the later stages of disease, along with episodes of "freezing"-complete arrest of gait when the patient is confronted by doorway or a narrow path between pieces of furniture. It becomes difficult for the patient to stand up from a seated position, or to turn over in bed. Impairment of fine motor control impairs activities of daily living such as fastening buttons, writing (micrographia), eating with knife and fork, shaving, and hair-combing. It becomes difficult to perform two activities simultaneously, such as walking and talking.

Tremor. Only about half of all PD patients have tremor early in the course of the disease; the rest usually develop it as the disease progresses. It is typically most pronounced in the hands (pill-rolling tremor) and is seen mainly when the affected limbs are at rest, improving or disappearing with voluntary movement. Its frequency is ca. 5 Hz, it is often asymmetrical, and it can be exacerbated by even mild stress (mental calculations, etc.).

Rigidity. Elevated muscle tone is felt by the patient as muscle tension or spasm and by the examiner as increased resistance to passive movement across the joints. Examination may reveal *cogwheel rigidity*, i.e., repeated, ratchet-like oscillations of resistance to passive movement across the wrist, elbow, or other joints, which may be brought out by alternating passive flexion and extension.

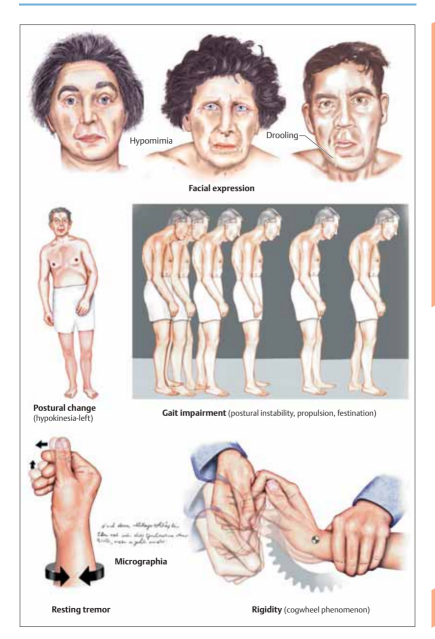
Postural instability (loss of balance). Propulsion and retropulsion arise in the early stages of Parkinson disease because of generalized impairment of the postural reflexes that maintain the bipedal stance. Related phenomena include involuntary acceleration of the gait (*festination*), difficulty in stopping walking, gait instability, and frequent falls.

Accompanying Manifestations

Behavioral Changes

Depression. The range of depressive manifestations includes worry, anxiety, avoidance of social contact, general unhappiness, listlessness, querulousness, brooding, somatoform disturbances, and (rarely) suicidal ideation.

Anxiety. Tension, worry, mental agitation, lack of concentration, and dizziness are relatively common complaints.



Dementia. Impairment of memory and concentration in early PD-associated dementia may be difficult to distinguish from depressive manifestations. The side effects of pharmacotherapy (p. 212) must be kept in mind before treatment is initiated for patients suffering from disorientation, confusion, suspiciousness, and other emotional changes. Impaired memory is usually not a major feature of PD; as the disease progresses, about 20% of patients develop decreased flexibility of thought and action, perseveration, and increasing difficulty in planning future activities. The development of dementia in PD is correlated with an increase in the number of Lewy bodies (pp. 210).

Hallucinations. A state of excessive suspiciousness, vivid dreams, and increasing anxiety may evolve into one of severe confusion with visual hallucinations. Frank psychosis (e.g., paranoid delusions, ideas of reference, or delusional jealousy) may be due to other causes than PD, particularly an adverse effect of antiparkinsonian medication. Dementia with Lewy bodies (a syndrome in which the clinical features of Parkinson disease are found together with dementia, fluctuating level of consciousness, visual hallucinations, and frequent falls) is another possible cause, especially in patients who are unusually sensitive to low doses of neuroleptics (⇒ exacerbation of parkinsonism, delirium, malignant neuroleptic syndrome, p. 347).

Autonomic Dysfunction

Blood pressure changes. Hypotension is a common side effect of antiparkinsonian medications (levodopa, dopamine agonists). Marked orthostatic hypotension, if present, suggests the possible diagnosis of multisystem atrophy.

Constipation may be caused by autonomic dysfunction, as a manifestation of the disease, or as a side effect of medication (anticholinergic agents).

Bladder disorders. Polyuria, urinary urgency, and urinary incontinence occur mainly at night and in patients with severe akinesia (who have difficulty getting to the toilet). PD only rarely causes severe bladder dysfunction.

Sleep disorders. PD commonly causes disturbances of the sleep–wake cycle, including difficulty falling asleep, nocturnal breathing prob-

lems similar to sleep apnea syndrome, and shortening of the sleep cycle. Sleep may also be interrupted by nocturnal akinesia, which makes it difficult for the patient to turn over in bed.

Sexual dysfunction. Spontaneous complaints of diminished libido or impotence are rare. Increased libido is a known side effect of levodopa and dopamine agonists.

Hyperhidrosis. Mainly occurs as generalized, irregular, sudden episodes of sweating.

Seborrhea. Mainly on the forehead, nose, and scalp (greasy face, seborrheic dermatitis).

Leg edema is often the result of physical inactivity.

Sensory Manifestations

Pain in the arm or shoulder, sometimes accompanied by fatigue and weakness, may be present for years before the cardinal manifestations arise and enable a diagnosis of PD. Back pain and nuchal cramps are frequent secondary effects of parkinsonian rigidity and abnormal posture. Dystonia may also come to attention because of the pain it produces.

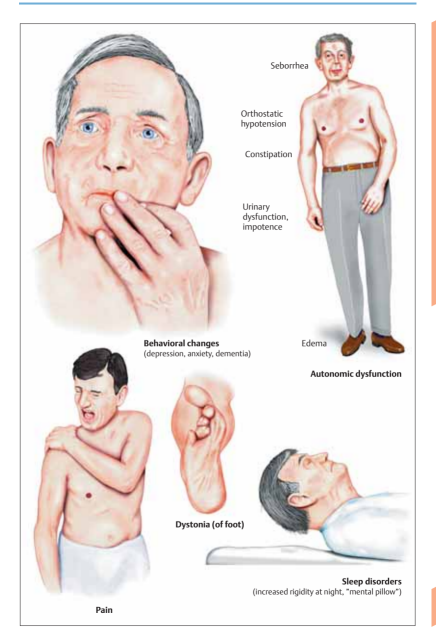
Dysesthesia. Heat, burning or cold sensations may be felt in various parts of the body. For *restless legs syndrome*, see p. 114.

Other Motor Manifestations

Dystonia. Tonic dorsiflexion of the big toe with extension or flexion of the other toes may occur in the early morning hours or during walking. Dystonia may be drug-induced (e.g., by levodopa) or due to the disease itself. The differential diagnosis includes dopa-responsive dystonia (a disorder of autosomal dominant inheritance) and Wilson disease (p. 307), two predominantly dystonic motor disorders with onset in childhood and adolescence.

Visual disturbances are caused by impairment of eye movement. Vertical gaze palsy is suggestive of *progressive supranuclear palsy* (p. 302). The reduced blinking rate of PD may lead to a burning sensation on the cornea, or to conjunctivitis.

Parkinson Disease: Clinical Features



Basal ganglia. The basal ganglia consist of the caudate nucleus (CN), putamen, globus pallidus (= pallidum; GPe = external segment, GPi = internal segment; putamen + pallidum = lentiform nucleus), claustrum, substantia nigra (SN; SNc = pars compacta, SNr = pars reticularis), and the subthalamic nucleus (STN). CN + putamen = (dorsal) striatum; nucleus accumbens + portions of olfactory tubercle + anterior portion of putamen + CN = limbic (ventral) striatum. Substantia nigra (SN): The SNr (ventral portion of SN) contains small amounts of dopamine and iron, giving it a reddish color, while the SNc (dorsal portion) contains large quantities of dopamine and melanin, making it black (whence the name, substantia nigra).

Connections. The basal ganglia are part of a number of parallel and largely distinct (segregated) neural pathways (circuits). Each circuit originates in a cortical area that is specialized for a specific function (skeletal motor, oculomotor, associative-cognitive, or emotional-motivational control), passes through several relay stations in the basal ganglia, and travels by way of the thalamus back to the cerebral cortex. Cortical projection fibers enter the basal ganglia at the striatum (input station) and exit from the GPi and SNr (output station). Input from the thalamus and brain stem also arrives at the striatum. Within the basal ganglia, there are two circuits subserving motor function, the so-called direct and indirect pathways. The direct pathway runs from the putamen to the GPi and SNr, while the indirect pathway takes the following trajectory: putamen \Rightarrow GPe \Rightarrow STN \Rightarrow GPi \Rightarrow SNr. The GPi and SNr project to the thalamus and brain stem.

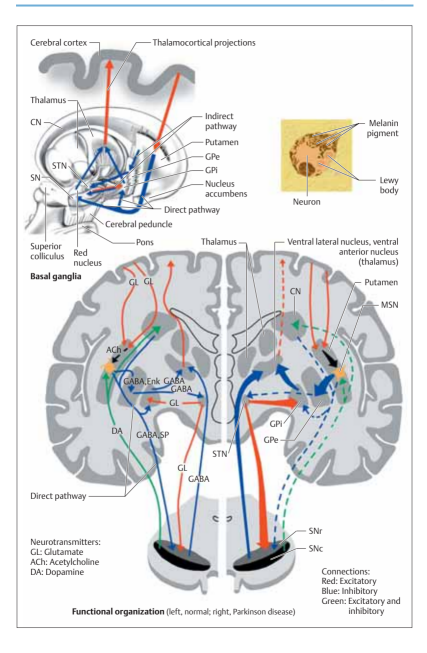
Neurotransmitters. *Glutamate* mediates excitatory impulses from the cortex, amygdala, and hippocampus to the striatum. Synapses from STN fibers onto cells of the GPi and SNr are also glutamatergic. Both the excitatory and the inhibitory projections of the SNc to the basal ganglia are dopaminergic. In the striatum, *dopamine* acts on neurons bearing D_1 and D_2 receptors, of which there are various subtypes (D_1 group: d_1 , d_5 ; D_2 group: d_2 , d_3 , d_4). D_1 receptors predominate in the direct pathway, D_2 receptors in the indirect pathway. Cholinergic interneurons in the striatum form a relay station within the basal ganglia (transmitter: *acetylcholine*). *Medium spiny-type*

neurons (MSN) in the striatum have inhibitory projections to the GPe, GPi, and SNr (transmitters: GABA, substance P/SP, enkephalin/Enk). Other inhibitory GABAergic projections run from the GPi to the STN, from the GPi to the thalamus (ventrolateral and ventroanterior nucleus), and from the SN to the thalamus. The thalamocortical projections are excitatory.

Motor function. The direct pathway is activated by cortical and dopaminergic projections to the striatum. The projection from the striatum in turn inhibits the GPi, diminishing its inhibitory output to the thalamic nuclei (i.e., causing net thalamic activation). Thalamocortical drive thus facilitates movement initiated in the cerebral cortex (voluntary movement). In the indirect pathway, the striatum, under the influence of afferent cortical and dopaminergic projections, exerts an inhibitory effect on the GPe and STN. The result is a diminished excitatory influence of the STN on the GPi and SNr, ultimately leading to facilitation of cortically initiated voluntary movement and inhibition of involuntary movement.

Pathophysiology

The cause of Parkinson disease is unknown. Its structural pathological correlate is a loss of neurons in the caudal and anterolateral parts of the SNc, with reactive gliosis and formation of Lewy bodies (eosinophilic intracytoplasmic inclusions in neurons) and Lewy neurites (abnormally phosphorylated neurofilaments) containing asynuclein. Loss of pigment in the substantia nigra can be seen macroscopically. The most prominent neurochemical abnormality is a deficiency of dopamine in the striatum, whose extent is directly correlated with the severity of PD. The physiological effect of the lack of (mostly inhibitory) dopamine neurotransmission in the striatum is a relative increase in striatal activity. in turn causing functional disinhibition of the subthalamic nucleus via the indirect pathway. Meanwhile, in the direct pathway, decreased striatal inhibition of the GPi enhances the inhibitory influence of the GPi on the thalamus. leading to reduced activity in the thalamocortical projection. These changes in neural activity manifest themselves in the clinically observable akinesia, rigidity, and postural instability. For tremor, see p. 62.



The *goal of treatment* is improvement of the motor, autonomic, and cognitive symptoms of the disease. The treatment generally consists of medication along with physical, occupational, and speech therapy. Neurosurgical procedures are mostly reserved for intractable cases (see below). Pharmacotherapy is palliative, not curative. It is begun when the patient has trouble carrying out the activities of daily living and is prescribed, not according to a uniform pattern, but in relation to the needs of the individual patient.

Symptomatic Treatment

Dopaminergic agents. Levodopa is actively absorbed in the small intestine and rapidly distributed throughout the body (especially to skeletal muscle). Amino acids compete with the levodopa transport system at the blood-brain barrier. A decarboxylase inhibitor that does not penetrate the blood-brain barrier (benserazide or carbidopa) is administered together with levodopa to prevent its rapid breakdown in the peripheral circulation. Once it reaches the brain, levodopa is decarboxylated to dopamine, which is used for neurotransmission in the striatum. After it has been released from the presynaptic terminals of dopaminergic neurons in the striatum and exerted its effect on the postsynaptic terminals, it is broken down by two separate enzyme systems (deamination by monoamine oxidase type B, MAO-B; methylation by catechol-O-methyltransferase, COMT). Levodopa effectively reduces akinesia and rigidity, but has only a mild effect against tremor. Its long-term use is often complicated by motor fluctuations, dyskinesia, and psychiatric disturbances. Dopamine agonists (DAs) mimic the function of dopamine, binding to dopamine receptors. Their interaction with D1 and D2 receptors is thought to improve motor function, while their interaction with D3 receptors is thought to improve cognition, motivation, and emotion. Long-term use of DAs is less likely to cause unwanted motor side effects than long-term use of levodopa. Commonly used DAs include bromocriptine (mainly a D₂ agonist), lisuride (mainly a D₂ agonist), and pergolide (a D₁, D₂, and D₃ agonist). Apomorphine, an effective D_1 and D_2 agonist, can be given by subcutaneous injection, but its effect lasts only about 1 hour. Other, recently introduced dopamine agonists are ropinirol and pramipexol (D_2 and D_3), cabergoline (D_2), and α -dihydroergocryptine (mainly D_2).

Selegiline inhibits MAO-B selectively and irreversibly (\Rightarrow reduced dopamine catabolism \Rightarrow increase in striatal dopamine concentration). *Entacapone* increases the bioavailability of levodopa via peripheral inhibition of COMT.

Nondopaminergic agents. Anticholinergic agents (biperidene, bornaprine, metixene, trihexyphenidyl) act on striatal cholinergic interneurons. Budipine can relieve tremor (risk of ventricular tachycardia ⇒ ECG monitoring). Glutamate antagonists (amantadine, memantine) counteract increased glutamatergic activity at the N-methyl-D-aspartate (NMDA) glutamate receptor in the indirect pathway.

Transplant Surgery

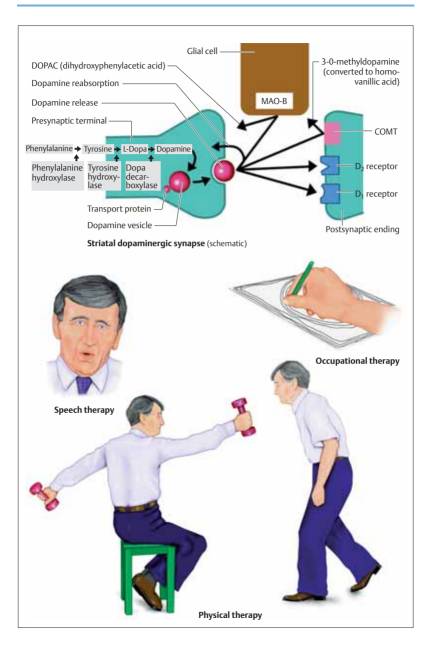
Current research on intrastriatal transplantation of stem cells (derived from fetal tissue, from umbilical cord blood, or from bone marrow) seems promising.

Stereotactic Neurosurgical Procedures, Deep Brain Stimulation (for abbreviations, see p. 210)

These procedures can be used when PD becomes refractory to medical treatment. *Pallidotomy* (placement of a destructive lesion in the GPi) derives its rationale from the observed hyperactivity of this structure in PD. *Deep brain stimulation* requires bilateral placement of stimulating electrodes in the GPi or STN. Highfrequency stimulation by means of a subcutaneously implanted impulse generator can improve rigor, tremor, akinesia, and dyskinesia.

Genetics of PD

A genetic predisposition for the development of PD has been postulated. Mutations in the genes for α -synuclein (AD), parkin (AR), and ubiquitin C-terminal hydrolase L1 (UCHL1; AD) have been found in pedigrees affected by the rare autosomal dominant (AD) and autosomal recessive (AR) familial forms of PD.



Multiple sclerosis (MS) is characterized by multiple symptoms and signs of brain and spinal cord dysfunction that are disseminated in both time and space. Its pathological hallmark is inflammatory demyelination and axonal lesions; its etiology remains unknown at present despite decades of intensive investigation.

A relapse is the appearance of a new neurological disturbance, or the reappearance of one previously present, lasting at least 24 hours. All such disturbances arising within a one-month period are counted as a single relapse. The relapse rate is the number of relapses per year. Clear improvement of neurological function is termed remission.

The course of MS varies greatly from one individual to another, but two basic types of course can be identified: relapsing-remitting (66-85%; most common when onset is before age 25; well-defined relapses separated by periods of nearly complete recovery with or without residual symptoms; does not progress during remission) and chronic progressive. The latter can be divided into three subtypes: primary chronic progressive (9-37%; most common when onset is after age 40; progresses from disease onset onward); secondary progressive (seen in over 50% of cases 6-10 years after onset; initially remitting-relapsing, later chronically progressive; recurrences, mild remissions, and plateau phases may occur); and progressively remitting-relapsing (rare; complete remission may or may not occur after relapses; symptoms tend to worsen from one relapse to the next).

Clinical Manifestations

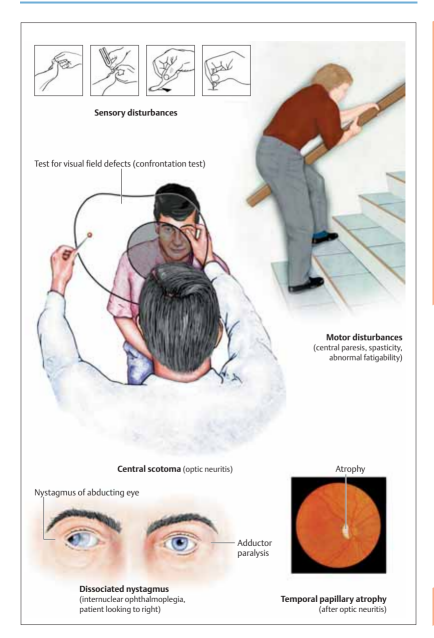
The symptoms and signs of MS reflect dysfunction of the particular areas of the nervous system involved and are not specific for this disease. Typical MS manifestations include paralysis, paresthesiae, optic neuritis (*retrobulbar neuritis*), diplopia, and bladder dysfunction.

Paresis, spasticity, fatigability. Upper-motorneuron type paralysis of the limbs either is present at onset or develops during the course of MS. Involvement is often asymmetrical and mainly in the legs, especially in the early stage of the disease. Spasticity makes its first appearance in the form of extensor spasms; flexor spasms develop later. The latter are often painful, cause frequent falls, and, if severe and persistent, can cause flexion contractures (*paraplegia in flexion*). Many patients complain of abnormal fatigability.

Sensory manifestations. Episodic or continuous paresthesiae (sensations of tingling or numbness, tightness of the skin, heat, cold, burning, prickling) are common, particularly in the early stage of the disease, with or without other manifestations of neurological dysfunction. As the disease progresses, such positive phenomena usually recede and are replaced by sensory deficits affecting all sensory modalities. A constant or only slowly rising sensory level ("sensory transverse cord syndrome") is uncharacteristic of MS and should prompt the search for a spinal cord lesion of another kind. Many MS patients have Lhermitte's sign (which is actually a symptom), an electric or coldlike paresthesia traveling from the nuchal region down the spine, sometimes as far as the legs, on flexion of the neck (p. 49). If no other symptoms or signs are present, other causes should be considered (e.g., a cervical spinal cord tumor).

Pain in MS most often appears in the form of trigeminal neuralgia (p. 186), severe pain in the limbs (p. 108), tonic spasms (p. 204), or backaches, sometimes with radiation in a radicular pattern. Other painful phenomena include flexor spasms due to spasticity, contractures, and dysuria due to urinary tract infection.

Visual impairment in MS is usually due to optic neuritis (mostly unilateral), which also produces pain in or around the eye. The impairment begins as blurred or clouded vision and progresses to cause reading impairment and visual field defects (central scotoma or diffuse defects). Marcus Gunn pupils (p. 92) may be observed. Physical exercise, high ambient temperature, menstruation, or cigarette smoking can aggravate existing visual problems (Uhthoff's phenomenon). Optic neuritis, as an isolated finding, is not necessarily the first manifestation of MS; patients with bilateral optic neuritis have a much lower risk of developing MS than those with the unilateral form. Diplopia is usually due to internuclear ophthalmoplegia (p. 86). Nystagmus (p. 88).



200

Multiple Sclerosis

Incoordination. Intention tremor, dysarthria, truncal ataxia, and oculomotor dysfunction are common. Gait unsteadiness due to motor incoordination is often experienced by the patient as dizziness or lightheadedness. Acute vertigo with nausea, vomiting, and nystagmus can also occur.

Autonomic dysfunction. Bladder dysfunction (p. 156) frequently develops in the course of MS, causing problems such as urinary urgency, incomplete voiding, or urinary incontinence. Urinary tract infection is a not infrequent result. Fecal incontinence (p. 154) is rare, but constipation is common. Sexual dysfunction (e.g., erectile dysfunction or loss of libido) is also common and may be aggravated by spasticity or sensory deficits in the genital region. Psychological factors such as depression, insecurity, and marital conflict often play a role as well. If its cause is organic, sexual dysfunction in MS is usually accompanied by bladder dysfunction.

Behavioral changes. Mental changes (depression, marital conflict, anxiety) and cognitive deficits of variable severity can occur both as a reaction to and as a result of the disease.

Paroxysmal phenomena in MS include epileptic seizures, trigeminal neuralgia, attacks of dysarthria with ataxia, tonic spasms, episodic dysesthesiae, pain, and facial myokymia.

Differential Diagnosis

There is no single clinical test, imaging study, or laboratory finding that alone establishes the diagnosis of MS (p. 218; Table 27, p. 375). A meticulous differential diagnostic evaluation is needed in every case.

(Cerebral) Vasculitis (p. 180). Systemic lupus erythematosus, Sjögren syndrome, Behçet syndrome, granulomatous angiitis, polyarteritis nodosa, antiphospholipid syndrome, chronic inflammatory demyelinating polyradiculoneuropathy (CIDP, p. 328).

Inflammatory diseases. Neurosarcoidosis, neuroborreliosis, neurosyphilis, Whipple disease, postinfectious acute disseminated encephalomyelitis (ADEM), progressive multifocal leukoencephalopathy (PML), subacute sclerosing panencephalitis (SSPE), HIV infection, HTLV-1 infection. **Neurovascular disorders.** Arteriovenous fistula of spinal dura mater, cavernoma, CADASIL (p. 172).

Hereditary/metabolic disorders. Spinocerebellar ataxias, adrenoleukodystrophy, endocrine diseases, mitochondrial encephalomyelopathy, vitamin B₁₂ deficiency (funicular myelosis).

Tumors of the brain or spinal cord (e.g., lymphoma, glioma, meningioma).

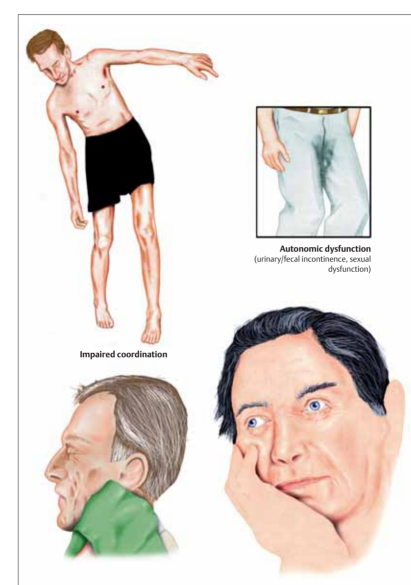
Skull base anomalies. Arnold–Chiari malformation, platybasia.

Myelopathy. Cervical myelopathy (spinal stenosis).

Somatoform disturbances in the context of mental illness.

Prognosis

Favorable prognostic indicators in MS include onset before age 40, monosymptomatic onset, absence of cerebellar involvement at onset. rapid resolution of the initial symptom(s), a relapsing-remitting course, short duration of relapses, and long-term preservation of the ability to walk. A relatively favorable course is also predicted if, after the first 5 years of illness, the MRI reveals no more than a few, small lesions without rapid radiological progression and the clinical manifestations of cerebellar disease and central paresis are no more than mild. A benign course, defined as a low frequency of recurrences and only mild disability in the first 15 years of illness, is seen in 20-30% of patients. The disease takes a malignant course, with major disability within 5 years, in fewer than 5% of patients. Half of all MS patients have a second relapse within 2 years of disease onset.



Paroxysmal symptoms (trigeminal neuralgia)

Behavioral changes

Diagnosis

Patients with MS are evaluated by clinical examination, laboratory testing, neuroimaging, and neurophysiological studies. The clinical manifestations of MS and the lesions that cause them vary over the course of the disease (*dissemination in time and space*). Diagnostic classification is problematic (p. 216) if only one lesion is found (e.g., by MRI), if symptoms and signs are in only one area of the CNS (e.g., spinal cord), or if only one attack has occurred (Table 27, p 375).

Clinical Manifestations

Sensory deficits, upper-motor-neuron paresis, incoordination, visual impairment (field defects), nystagmus, internuclear ophthalmoplegia, and/or bladder dysfunction are common signs of MS. Complaints of pain, paresthesiae, abnormal fatigability, or episodic disturbances are often, by their nature, difficult to objectify. Clinical examination may reveal no abnormality because of the episodic nature of the disease itself.

Laboratory Tests and Special Studies

Evoked potentials. Visual evoked potential (VEP) studies reliably detect optic nerve lesions, but neuroimaging is better for detecting lesions of the optic tract or optic radiation. VEP reveals prolongation of the P100 latency in one eye and/ or an abnormally large discrepancy between the latencies in the two eyes in roughly 40% of MS patients without known optic neuritis, and in almost half of those with early optic neuritis. Somatosensory evoked potential (SEP) studies of the median or tibial nerve typically reveal prolonged latencies in MS. Low amplitude of evoked potentials, on the other hand, often indicates a pathological process of another type, e.g. tumor. SEP abnormalities are found in up to 60% of MS patients with predominantly sensory manifestations. Auditory evoked potential (AEP) studies are less sensitive in MS than VEP or SEP. The most common AEP change is prolongation of latency. AEP studies are helpful for the further classification of vertigo, tinnitus, and hearing loss. Motor evoked potential (MEP) studies reveal prolonged central conduction times when CNS lesions involve the pyramidal pathway. The

sensitivity of MEP in MS is approximately the same as that of SEP. MEP studies can provide supporting evidence for MS in patients with latent paresis, gait disturbances, abnormal reflexes, or movement disorders that are difficult to classify.

Tests of bladder function. The residual urine volume can be measured by ultrasound. It should not exceed 100 ml in patients with a normal bladder capacity of 400–450 ml; in general, it should normally be 15–20% of the cystomanometrically determined bladder volume. Urodynamic electromyography (EMG) provides more specific data concerning bladder dysfunction.

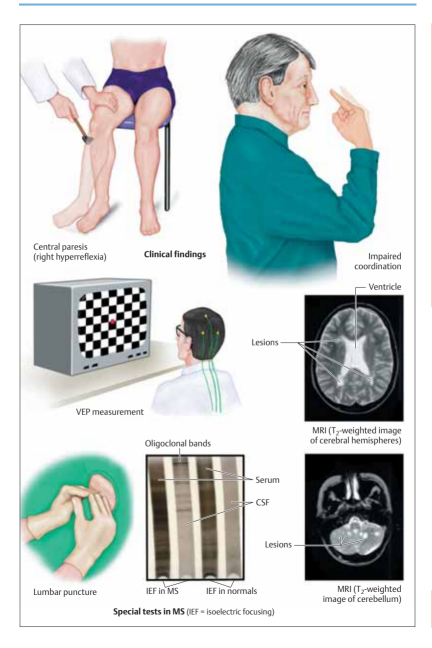
Neuroimaging. CT may reveal other diseases that enter into the differential diagnosis of MS (e. g., brain tumor) but is insufficiently sensitive (ca. 25–50%) to be useful in diagnosing MS itself. MRI scans reveal the characteristic foci of demyelination disseminated in the CNS (MS plaques); contrast enhancement is seen in acute but not in chronic lesions. The sensitivity of MRI for MS is greater than 90%, but its specificity is considerably lower; thus, the MRI findings alone cannot establish the diagnosis.

CSF examination. CSF abnormalities are found in more than 95% of MS patients. The cell count rarely exceeds 20 cells/mm³. The total protein concentration is elevated in ca. 40% of patients, and intrathecal IgG synthesis (IgG index) in ca. 90%. Oligoclonal IgG is found in 95% of MS patients, and antibodies to mumps, measles and herpes zoster in 80%.

Pathogenesis

The *early* course of MS varies among patients in accordance with the variable extent of the inflammatory lesions and disturbances of the blood–brain-barrier. The severity of *late* manifestations is correlated with the number of plaques. It is hypothesized that MS is caused by a combination of genetic (polygenic) predisposition and exogenous factors (viral or bacterial infection?) that induces an inappropriate immune response to one or more CNS autoantigens (see p. 220) that have not yet been identified.

Multiple Sclerosis



Activation. Circulating autoreactive CD4⁺ T lymphocytes bear antigen-specific surface receptors and can cross the blood-brain barrier (BBB) when activated, e.g., by neurotropic viruses, bacterial superantigens, or cytokines. In MS, activated T lymphocytes react with MBP, PLP, MOG, and MAG. Circulating antibodies to various components of myelin can also be detected (for abbreviations, see below¹).

Passage through the BBB. Activated lymphocytes and myelinotoxic antibodies penetrate the BBB at the venules (perivenous distribution of inflammation).

Antigen presentation and stimulation. In the CNS, antigen-presenting cells (microglia), recognition molecules (MHC class II antigens), and co-stimulatory signals (CD28, B-7.1) trigger the renewed activation and clonal proliferation of incoming CD4+ T lymphocytes into T_H1 and T_H2 cells. Proinflammatory cytokines elaborated by the T_H1 cells (IL-2, IFN- γ , TNF- α , LT)² induce phagocytosis by macrophages and microglia as well as the synthesis of mediators of inflammation (TNF- α , OH-, NO)² and complement factors. The T_H2 cells secrete cytokines (IL-4, IL-5, IL-6)² that activate B cells (⇒ myelinotoxic autoantibodies, complement activation), ultimately causing damage to myelin. The T_H2 cells also produce IL-4 and IL-10, which suppress the T_H1 cells.

Demyelination. Lesions develop in myelin sheaths (which are extensions of oligodendroglial cell membranes) and in axons when the inflammatory process outstrips the capacity of repair mechanisms.

Scar formation. The inflammatory response subsides and remyelination of damaged axons begins once the autoreactive T cells die (apoptosis), the BBB is repaired, and local anti-inflammatory mediators and cells are synthesized. Astroglia form scar tissue that takes the place of the dead cells. Axonal damage seems to be the

¹MBP, myelin basic protein; MOG, myelin-oligodendrocyte glycoprotein; MAG, myelin-associated glycoprotein; PLP, proteolipid protein; S100 protein, CNPase, αβ-crystallin, transaldolase

²IL, interleukin; IFN- γ , interferon-gamma; TNF- α , tumor necrosis factor-alpha; LT, lymphotoxin; OH-: hydroxyl radical; NO, nitric oxide

³p.o., orally; s.c., subcutaneously; i.m., intramuscularly; i. v., intravenously.

main cause of permanent neurological deficits, as dystrophic axons apparently cannot be remyelinated.

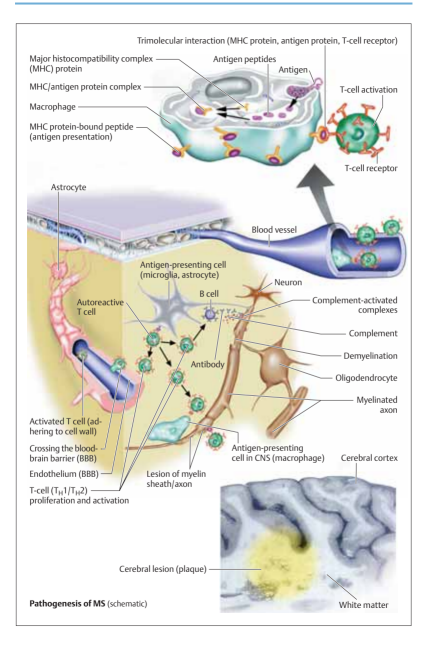
Treatment

Relapse is treated with high-dose corticosteroids, e. g., methylprednisolone, 1 g/day for 3–5 days, which produce (unselective) immunosuppression, reduce BBB penetration by T cells, and lessen T_H1 cytokine formation. Plasmapheresis may be indicated in refractory cases. **Drugs that reduce the frequency and intensity of relapses.** *Azathioprine* p.o. (immunosuppression via reduction of T cell count), interferon beta-1b and beta-1a s.c. or i.m. (cytokine modulation, alteration of T-cell activity), glatiramer acetate s.c. (copolymer-1; blocks/competes at binding sites for encephalitogenic peptides on MHC-II molecules), IgG i. v. (multiple modes of action), and natalizumab (selective adhesion inhibitor).

Drugs that delay secondary progression. Interferon beta-1b and beta-1a. *Mitoxantrone* suppresses B cells and decreases the CD4/CD8 ratio. *Methotrexate* and *cyclophosphamide* (different dosage schedules) delay MS progression mainly by unselective immunosuppression and reduction of the T-cell count.

Slowing of primary progression. No specific therapy is known at present.

Symptomatic therapy/rehabilitation. Medications, physical, occupational, and speech therapy, social, psychological, and dietary counseling, and mechanical aids (e.g., walking aids, wheelchair) are provided as needed. The possible benefits of oligodendrocyte precursor cell transplantation for remyelination, and of growth factors and immunoglobulins for the promotion of endogenous remyelination, are currently under investigation in both experimental and clinical studies.



Syndromes

Localization

CNS infection may involve the leptomeninges and CSF spaces (*meningitis*), the ventricular system (*ventriculitis*), the gray and white matter of the brain (*encephalitis*), or the spinal cord (*myelitis*). A focus of bacterial infection of the brain is called a *brain abscess*, or *cerebritis* in the early stage before a frank abscess is formed. Pus located between the dura mater and the arachnoid membrane is called a *subdural empyema*, while pus outside the dura is called an *epidural abscess*.

Course

The clinical manifestations may be *acute* (purulent meningitis, CNS listeriosis, herpes simplex encephalitis), *subacute* (cerebral abscess, focal encephalitis, neuroborreliosis, neurosyphilis, tuberculous meningitis, actinomycosis, nocardiosis, rickettsiosis, neurobrucellosis), or *chronic* (tuberculous meningitis, neurobyphilis, neuroborreliosis, Whipple encephalitis, Creutzfeldt–Jakob disease). The epidemiological pattern of infection may be *sporadic, endemic* or *epidemic*, depending on the pathogen.

Clinical Manifestations

Meningitis and encephalitis rarely occur as entirely distinct syndromes; they usually present in mixed form (meningoencephalitis, encephalomyelitis). CSF examination establishes the diagnosis.

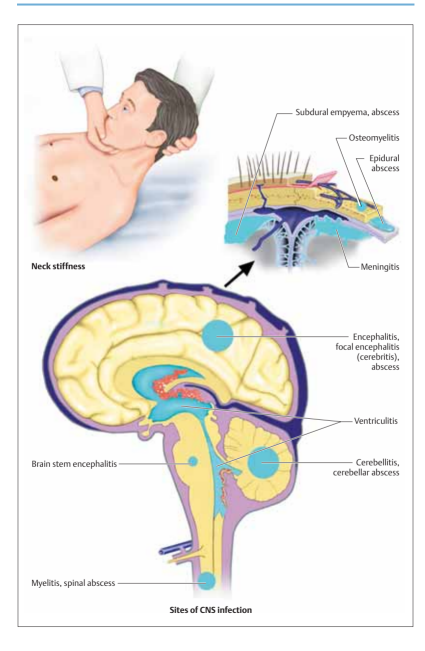
These disorders may present in specific ways in certain patient groups. *Neonates and children* commonly manifest failure to thrive, fever or hypothermia, restlessness, breathing disorders, epileptic seizures, and a bulging fontanelle. *The elderly* may lack fever but frequently have behavioral abnormalities, confusion, epileptic seizures, generalized weakness, and impairment of consciousness ranging to coma. *Immunodeficient patients* commonly have fever, headache, stiff neck, and drowsiness in addition to the manifestations of their primary illness.

Meningitic syndrome is characterized by fever, severe, intractable headache and backache, photophobia and phonophobia, nausea, vomiting, impairment of consciousness, stiff neck, and hyperextended posture, with opisthotonus or neck pain on flexion. Kernig's sign (resistance to passive raising of leg with extended knee) and Brudzinski's sign (involuntary leg flexion on passive flexion of the neck) are signs of meningeal involvement. Painful neck stiffness is due to (lepto)meningeal irritation infectious bv meningitis, septicemia, subarachnoid hemorrhage, neoplastic meningitis, or other causes. Isolated neck stiffness not caused by meningitis (meningism) may be due to cervical disorders such as arthrosis, fracture, intervertebral disk herniation, tumor, or extrapyramidal rigidity. Papilledema is usually absent; when present, it indicates intracranial hypertension (p. 158).

Encephalitic syndrome is characterized by headache and fever, sometimes accompanied by epileptic seizures (often focal), focal signs (cranial nerves deficits, especially of CN III, IV, VI, and VII; aphasia, hemiparesis, hemianopsia, ataxia, choreoathetosis), behavioral changes, and impairment of consciousness (restlessness, irritability, confusion, lethargy, drowsiness, coma). The neurological signs may be preceded by limb pain (myalgia, arthralgia), a slight increase in body temperature, and malaise. For *acute cerebellitis* (⇔ ataxia), see p. 276. *Brain stem encephalitis* produces ophthalmoplegia, facial paresis, dysarthria, dysphagia, ataxia, and hearing loss.

Myelitic syndrome. Myelitis presents with severe local pain, paraparesis, paresthesiae, or some combination of these. Incomplete or complete paraplegia or quadriplegia (p.48) develops within a few hours (acute) or days (subacute). The differential diagnosis may be difficult.

CNS Infections



Pathogenesis

Pathogens usually reach the CNS by local extension from a nearby infectious focus (e.g. sinusitis, mastoiditis) or by hematogenous spread from a distant focus. The ability of pathogens to spread by way of the bloodstream depends on their virulence and on the immune status of the host. They use special mechanisms to cross or circumvent the blood-brain barrier (p. 8). Some pathogens enter the CNS by centripetal travel along peripheral nerves (herpes simplex virus type I, varicella-zoster virus, rabies virus), others by endocytosis (Neisseria meningitidis), intracellular transport (Plasmodium falciparum via erythrocytes, Toxoplasma gondii via macrophages), or intracellular invasion (Haemophilus influenzae). Those that enter the subarachnoid space probably do so by way of the choroid plexus, venous sinuses, or cribriform plate (p. 76). Having entered the CSF spaces, pathogens trigger an inflammatory response characterized by the release of complement factors and cytokines, the influx of leukocytes and macrophages, and the activation of microglia and astrocytes. Disruption of the blood-brain barrier results in an influx of fluids and proteins across the vascular endothelium and into the CNS, causing vasogenic cerebral edema (p. 162), which is accompanied by both cytotoxic cellular edema and interstitial edema due to impaired CSF circulation. Cerebral edema causes intracranial hypertension. These processes, in conjunction with vasculitis, impairment of vascular autoregulatory mechanisms, and/or fluctuations of systemic blood pressure, lead to the development of ischemic, metabolic, and hypoxic cerebral lesions (focal necrosis, territorial infarction).

Treatment (Table 28, p. 375)

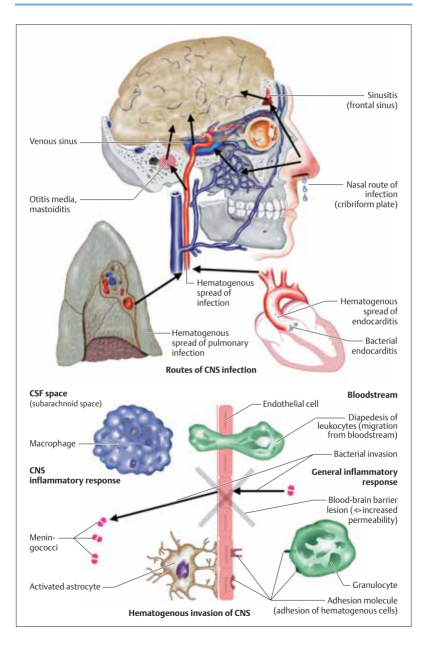
The immune system is generally no longer able to hold pathogens in check once they have spread to the CNS, as the immune response in the subarachnoid space and the neural tissue itself is less effective than elsewhere in the body. Having gained access to the CNS, pathogens meet with favorable conditions for further spread within it.

Prophylaxis. The occurrence and spread of CNS infection can be prevented by *mandatory re*-

porting (as specified by local law), prevention of exposure (isolation of sources of infection, disinfection, sterilization), and prophylaxis in persons at risk (active and passive immunization, chemoprophylaxis).

Treatment. Patients with bacterial or viral meningoencephalitis must be treated at once. The treatment strategy is initially based on the clinical and additional findings. Antimicrobial therapy is first given empirically in a broad-spectrum combination, then specifically tailored in accordance with the species and drug sensitivity pattern of the pathogen(s) identified. Causative organisms may be found in the CSF, blood, or other bodily fluids (e.g., throat smear, urine or stool samples, bronchial secretions, gastric juice, abscess aspirate).

CNS Infections



Bacterial Infections

Meningitis/Meningoencephalitis

For an overview of the most common pathogens, cf. Table 29 (p 376). *Immune prophylaxis:* Vaccines are available against *Haemophilus influenzae* type B infection (for infants, small children, and children over 6 years of age at increased risk), *Pneumococcus* (children over 2 years of age and adults with risk factors such as immunosuppression or asplenia), and *meningococcus* (travel to endemic regions, local outbreaks). *Chemoprophylaxis* is indicated for close contacts of persons infected with *Haemophilus influenzae* (rifampicin) or meningococcus (rifampicin, ciprofloxacin, or ceftriaxone).

Brain Abscess

Brain abscess begins as local cerebritis and is then transformed into an encapsulated region of purulent necrosis with perifocal edema. The pathogenic organisms may reach the brain by local or hematogenous spread (mastoiditis, otitis media, sinusitis, osteomyelitis; endocarditis, pneumonia, tooth infection, osteomyelitis, diverticulitis), or by direct inoculation (trauma, neurosurgery). The clinical manifestations include headache, nausea, vomiting, fever, impairment of consciousness, and focal or generalized epileptic seizures, neck stiffness, and focal neurological signs. The diagnosis is made by MRI and/or CT (which should include bone windows, as the infection may have originated in bony structures) and confirmed by culture of the pathogenic organism.

■ Bacterial Vasculitis (p. 180)

Arteries. Vessel wall inflammation in association with sepsis. Bacterial endocarditis causes cerebral abscess formation or infarction by way of infectious thromboembolism (⇔ focal inflammatory changes in the cerebral parenchyma ⇔ metastatic or embolic focal encephalitis). The syndrome is characterized by headache, fever, epileptic seizures, and behavioral changes in addition to focal neurological signs. *Meningoencephalitis* may cause arteritis by direct involvement of the vessels. Embolization of infectious material may lead to the development of *septic* ("mycotic") *aneurysms*. Veins. Bacterial *thrombophlebitis* of the cerebral veins or venous sinuses may arise as a complication of meningitis or by local spread of infection from neighboring structures.

Ventriculitis

Infection of the ventricular system (perhaps in connection with an intraventricular catheter for internal or external CSF drainage). The clinical findings are often nonspecific (somnolence, impairment of concentration and memory). Abdominal complaints (peritonitis) may predominate if the infection has spread down a ventriculoperitoneal shunt to the abdomen. *Diagnosis:* CSF examination and culture.

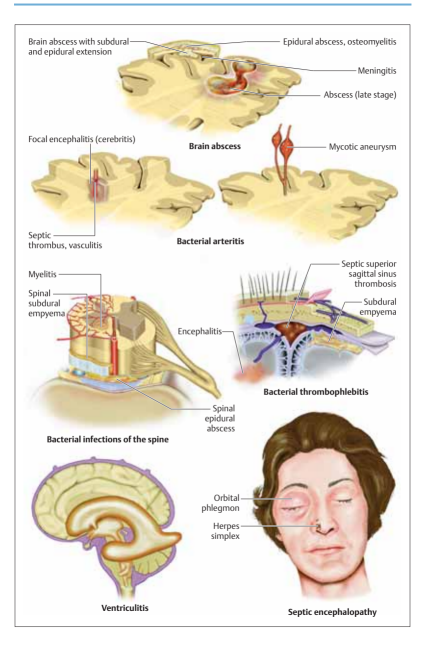
Septic Encephalopathy

Bacteremia leads to the release of endotoxins, which, in turn, impair cerebral function. Septic encephalopathy can produce findings suggestive of meningoencephalitis such as impairment of consciousness, epileptic seizures, paresis, and meningismus, despite the absence of CSF inflammatory changes and a sterile CSF culture. *Diagnosis*: EEG changes consistent with the diagnosis (general changes, triphasic complexes, burst suppression) in the setting of known systemic sepsis with sterile CSF. CT and MRI are normal.

226

Central Nervous System

CNS Infections



Lyme Disease (Neuroborreliosis)

Pathogenesis. The spirochete *Borrelia burg-dorferi* sensu lato (Europe: *B. garinii, B. afzelii*; North America: *B. burgdorferi* sensu stricto) is transmitted to man by ticks (Europe: *Ixodes ricinus*; North America: *Ixodes pacificus, I. scapularis*). The probability of infection is low unless the infected tick remains attached to the skin for at least 24–48 hours. Only 1–2% of individuals bitten by ticks become infected. The incubation time ranges from 3–30 days. The disease occurs in three stages, as described below.

Clinical manifestations. Stage I (localized infection). Up to 90% of all patients develop a painless, erythematous macule or papule that gradually spreads outward from the site of the tick bite in a ringlike or homogeneous fashion (erythema chronicum migrans). This is commonly accompanied by symptoms due to hematogenous spread of the pathogen, such as fever, fatigue, arthralgia, myalgia, or other types of pain, which may be the chief complaint, rather than the skin rash. Regional or generalized lymphadenopathy (lymphadenosis benigna cutis) is a less common presentation. All of these findings may resolve spontaneously.

Stage II (disseminated infection). Generalized symptoms such as fatigue, anorexia, muscle and joint pain, and headache develop in 10-15% of patients within ca. 3-6 weeks, sometimes accompanied by mild fever and neck stiffness. Cardiac manifestations: Myocarditis or pericarditis with AV block. Neurological manifestations: Cranial nerve palsies, painful polyradiculitis and lymphocytic meningitis (Bannwarth syndrome, meningopolyneuritis) are commonly seen in combination. One or more cranial nerves may be affected; the most common finding is unilateral or bilateral facial palsy of peripheral type. Neuroborreliosis-related polyradiculoneuropathy (which may be mistaken for lumbar disk herniation) is characterized by intense pain in a radicular distribution, most severe at night, with accompanying neurological deficits (motor. sensory, and reflex abnormalities, focal muscle atrophy). Borrelia-related meningitis (Lyme meningitis) usually causes alternating headache and neck pain, but the headache is mild or absent in some cases. It may be worst at certain times of day. CSF studies reveal a mononuclear

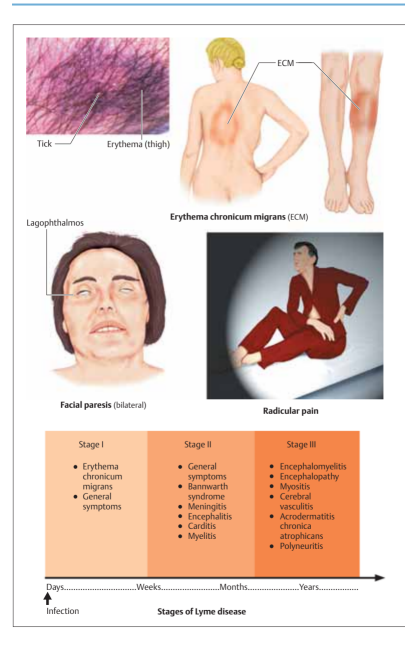
pleocytosis with a high plasma cell count and an elevated protein concentration, while the glucose concentration is normal. *Encephalitis* occurs relatively rarely and may cause focal neurological deficits as well as behavioral changes (impaired concentration, personality changes, depression). MRI reveals cerebral white-matter lesions, and the CSF findings are consistent with meningitis. *Myelitis*, when it occurs, often affects the spinal cord at the level of a radicular lesion.

Stage III (persistent infection). The latency from clinical presentation to the onset of stage III disease varies from 1 to 17 years (chronic Lyme neuroborreliosis). Few patients ever reach this stage, characterized by neurological deficits such as ataxia, cranial nerve palsies, paraparesis or quadriparesis, and bladder dysfunction (Lyme encephalomyelitis). Encephalopathy causing impairment of concentration and memory, insomnia, fatigue, personality changes, and depression has also been described. Myositis and cerebral vasculitis may also occur. In stage III of Lyme disease, acrodermatitis chronica atrophicans of the extensor surface of the limbs may be seen along with a type of polyneuropathy specific to Borrelia afzelii.

Diagnosis. Many patients have no memory of a tick bite. The diagnosis of Lyme disease is based on the presence of erythema chronicum migrans, the immunological confirmation of *Borrelia* infection (e.g., by ELISA, indirect immunofluorescence assay, Western blot, or specific IgG antibody–CSF-serum index) and/or the identification of the causative organism (e.g., by culture, histology, or polymerase chain reaction). By definition, the diagnosis also requires the presence of lymphocytic meningitis (with or without cranial nerve involvement or painful polyradiculoneuritis), encephalomyelitis, or encephalopathy.

Treatment. *Local symptoms:* Antibiotic such as doxycycline or amoxicillin (p.o.) for 3 weeks. *Neuroborreliosis:* Ceftriaxone or cefotaxime (i. v.) for 2–3 weeks. A vaccine has been approved for use in the United States, and another is being developed for use in Europe.

CNS Infections



Neurosyphilis

Pathogenesis. Syphilis is caused by the spirochete (bacterium) Treponema pallidum (TP) ssp. pallidum and is transmitted by direct exposure to infected lesions, usually on the skin or mucous membranes, during sexual contact. Other routes of transmission, such as the sharing of needles by intravenous drug users, are much less common. The disease has three clinical stages. In the primary and secondary stages, nonspecific tests (VDRL and RPR) and specific tests (TPHA, FTA-ABS, and 19S-(IgM-)FTA-ABS tests) yield positive results. Tertiary stage (currently rare): After an asymptomatic period of a few months to years (latent syphilis), organ manifestations develop, such as gummata (skin, bone, kidney, liver) and cardiovascular lesions (aortic aneurysm). The first year of the tertiary stage is designated the early latency period and is characterized by a high likelihood of recurrence and, thus, recurrent infectivity.

Clinical manifestations. TP may invade the nervous system at any stage of syphilis without necessarily producing signs or symptoms.

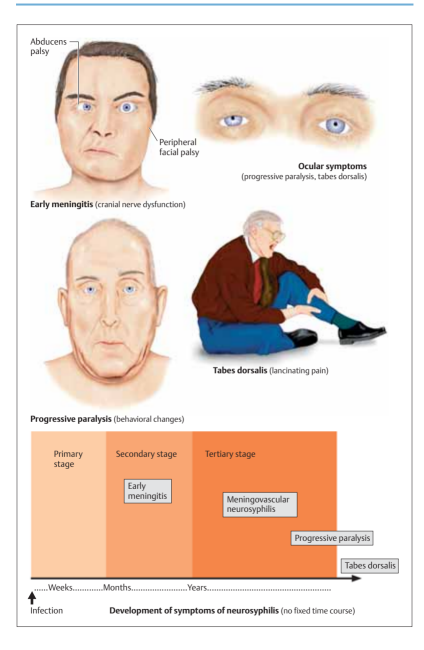
Early meningitis. A variably severe meningitic syndrome may be accompanied by deficits of CN VIII (sudden hearing loss), VII (facial palsy), or II (visual impairment). Meningopolyradiculitis is rare. CSF examination reveals lymphocytic pleocytosis (up to 400 cells/µl) and an elevated protein concentration. Meningitis resolves spontaneously, but late complications may occur. *Asymptomatic meningitis* (CSF changes in the absence of a meningitic syndrome) occurs in 20–30% of all infected persons.

Meningovascular neurosyphilis. Fluctuating symptoms such as headache, visual disturbances, and vertigo occur 5-12 years after the initial infection. Vasculitis (von Heubner angiitis) causes stroke, particularly in the territory of the middle cerebral artery, and may also affect small perforating vessels as well as cranial nerves (VIII, VII, V). Hydrocephalus, personality changes, epileptic seizures, and spinal cord signs (paraparesis, bladder dysfunction, anterior cord syndrome) round out the kaleidoscopic clinical picture. Gummata are rarely seen. CT and MRI findings suggest the diagnosis, and CSF examination reveals a mononuclear pleocytosis (up to 100 cells/µl), elevated protein concentration, elevated oligoclonal IgG, and VDRL positivity (up to 80%).

Progressive paralysis. Chronic meningoencephalitis with progressive paralysis occurs 10–25 years after the initial infection. The "preparalytic" stage, characterized by personality changes and mild impairment of concentration and memory, later evolves into the "paralytic" stage, characterized by more severe cognitive changes, dysarthria, dysphasia, tremor (mimic tremor), apraxia, gait impairment, urinary incontinence, and abnormal pupillary reflexes (roughly 25% of patients have Argyll–Robertson pupils, p. 92). The CSF findings resemble those of meningovascular syphilis.

Tabes dorsalis. This late meningovascular complication (25–30 years after the initial infection) produces ocular manifestations (Argyll–Robertson pupils, strabismus, papillary atrophy), pain (lightning pains = lancinating pain mainly in the legs; colicky abdominal pain), gait impairment (due to loss of acrognosis and proprioception), and autonomic dysfunction (impotence, urinary dysfunction). Joint deformities (Charcot joints) in the lower limbs are occasionally seen. The CSF cell count is relatively low, as in meningovascular syphilis.

Antibiotic therapy. The efficacy of treatment depends on the stage of disease in which it is instituted (the earlier, the better). Penicillin is the agent of choice.



Tuberculous Meningitis

Pathogenesis. **Mvcobacterium** tuberculosis transmission in man is usually by transfer of droplets from and to the respiratory tract (rarely orally or through skin lesions). The pathogen replicates in the lungs (primary infection), either in the lung tissue itself or within alveolar macrophages. Macrophages can only destroy tubercle bacilli after they have been activated by T cells: the course of the infection thus depends on the state of the immune system, i.e., on the ability of activated macrophages to hold the bacilli in check. The stage of primary infection lasts 2-4 weeks, is not necessarily symptomatic (if it is, then with nonspecific symptoms such as fever, anorexia, and lethargy), and cannot be detected by immune tests performed on the skin. The inflammatory process may also involve the regional (hilar) lymph nodes (primary complex). Calcified foci in the primary complex are easily seen on plain radiographs of the chest. The bacilli may remain dormant for years or may be reactivated when the patient's immune defenses are lowered by HIV infection, alcoholism, diabetes mellitus, corticosteroid therapy, or other factors (reactivated tuberculosis). Spread from the primary focus to other organs (organ tuberculosis) can occur during primary infection in immunocompromised patients, but only after reactivation in other patients. The bacilli presumably reach the CNS by hematogenous dissemination; local extension to the CNS from tuberculous bone (spinal cord, base of skull) is rare.

Symptoms and signs. The type and focus of CNS involvement (*neurotuberculosis*) vary, depending mainly on the age and immune status of the host.

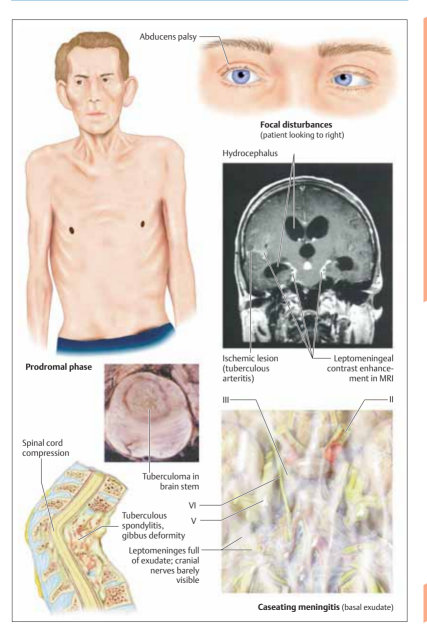
Tuberculous meningoecephalitis. The prodromal stage lasts 2–3 weeks and is characterized by behavioral changes (apathy, depression, irritability, confusion, delirium, lack of concentration), anorexia, weight loss, malaise, nausea, and fever. Headache and neck stiffness reflect *meningeal involvement*. Finally, *cerebral involvement* manifests itself in *focal signs* (deficits of CN II, III, VI, VII, and VIII; aphasia, apraxia, central paresis, focal epileptic seizures, SIADH) and/or *general signs* (signs of intracranial hypertension, hydrocephalus). The focal signs are caused by leptomeningeal adhesions, cerebral ischemia due to vasculitis, or mass lesions (tuberculoma). Chronic meningitis most likely reflects inadequate treatment, or resistance of the pathogen, rather than being a distinct form of the disease. Diagnosis: CSF examination for initial diagnosis and monitoring of disease course. The diagnosis of tuberculous meningitis can only be confirmed by detection of mycobacteria in the CSF with direct microscopic visualization, culture, or molecular biological techniques. As the prognosis of untreated tuberculous meningitis is poor, treatment for presumed disease should be initiated as soon as the diagnosis is suspected from the clinical examination and CSF findings; the latter typically include high concentrations of protein (several grams/liter) and lactate, a low glucose concentration (< 50% of blood glucose), a high cell count (over several hundred), and a mixed pleocytosis (lymphocytes, monocytes, granulocytes).

Tuberculoma is a tumorlike mass with a caseous or calcified core surrounded by granulation tissue (giant cells, lymphocytes). Tuberculomas may be solitary or multiple and are to be differentiated from tuberculous abscesses, which are full of mycobacteria and lack the surrounding granulation tissue. *Diagnosis*: CT or MRI.

Spinal tuberculosis. Transverse spinal cord syndrome can arise because of tuberculous myelomeningoradiculitis, epidural tuberculous abscess associated with tuberculous spondylitis/ discitis, or tuberculoma. *Diagnosis*: MRI.

Antibiotic treatment. One treatment protocol specifies a combination of isoniazid (with vitamin B_6), rifampicin (initially i.v., then p.o.), and pyrazinamide (p.o.). After 3 months, pyrazinamide is discontinued, and treatment with isoniazid and rifampicin is continued for a further 6–9 months. The treatment for HIV-positive patients includes up to five different antibiotics.

CNS Infections



Viral Infections

Viral Meningoencephalitis

Aseptic meningitis is characterized by a meningitic syndrome (p. 222) that arises acutely and takes a benign course over the ensuing 1 or 2 weeks, in the absence of signs of generalized or local infection (otitis media, craniospinal abscess, sinusitis), CSF findings: A mild granulocytic pleocytosis is seen in the first 48 hours and is then transformed into a mild lymphomonocvtic pleocytosis which can persist as long as 2 months after the clinical findings have regressed. The CSF protein and lactate concentrations are normal or only slightly elevated, while the CSF glucose concentration is normal or mildly decreased. The term "viral meningitis" is often used synonymously with aseptic meningitis, though, strictly speaking, the clinical picture of aseptic meningitis can also be produced by fungal, parasitic, or even bacterial infection (e.g., mycobacteria, mycoplasma, Brucella, spirochetes, Listeria, rickettsiae, incompletely treated bacterial meningitis). Aseptic meningitis may be postinfectious (HIV, rubella, measles, zoster) or postvaccinial or a sequela of sarcoidosis. Behcet disease, Vogt-Kovanagi-Harada svndrome. Mollaret meningitis, connective tissue diseases, and other, noninflammatory disorders (meningeal carcinomatosis, contrast agents, medications, subarachnoid hemorrhage, lead poisoning).

Pathogens. Their seasonal peak frequency is shown in the following table.

Viral meningitis. Frontal or retro-orbital headache, fever, and low-grade neck stiffness usually begin acutely and last for 1–2 weeks. The CSF findings are those of aseptic meningitis, described above; the IgG index or oligoclonal IgG may be elevated.

Viral encephalitis. The meninges are usually involved concomitantly (meningoencephalitis). Acute encephalitis mainly affects the gray matter and perivascular areas of the brain. Behavioral changes, psychomotor agitation, and focal epileptic seizures may be the leading symptoms (p. 192). The CSF findings are generally as listed above for aseptic meningitis, though pleocytosis may be absent at first. Diffuse and focal EEG changes are usually seen. CT and MRI often reveal pathological changes.

Acute demyelinating encephalomyelitis (ADEM) predominantly affects perivenous regions and the cerebral white matter (*leukoencephalitis*). The disease takes a variable course (a monophasic course with complete resolution is among the possibilities). ADEM can occur during or after a bout of infectious disease (measles, chickenpox, rubella, influenza) or after a vaccination (smallpox, measles, mumps, polio). Both encephalitic and myelitic syndromes can occur (spastic paraparesis or quadriparesis). The white-matter lesions are demonstrated best by MRI, less well by CT.

Summer, Early Spring	Autumn, Winter	Winter, Spring	All Year Round
Arboviruses, ente- roviruses	Lymphocytic chorio- meningitis virus	Mumps	HIV, herpes simplex virus, cytomegalovirus

The viral pathogens that most commonly cause meningitis differ from those most commonly causing encephalitis and myelitis (cf. Table 30, p. 376).

Identification of pathogen: Serologic tests or isolation of the virus from *throat smears* (poliovirus, coxsackievirus, mumps virus; ade-

novirus, HSV type 1), stool samples (coxsackievirus, polio virus), CSF (coxsackievirus, mumps, adenovirus, arbovirus, rabies, VZV, LCMV, HSV type 2), blood (arbovirus, EBV, LCMV, CMV, HSV type 2), urine (mumps, CMV), or saliva (mumps, rabies).



Personality change (perseveration, apraxia, aphasia)



(hallucinations, psychomotor hyperactivity, loss of coordination, fluctuating level of consciousness)



Extrapyramidal motor dysfunction (tonic upward gaze deviation)

Focal signs (partial epileptic seizure)

Loss of drive (anxiety, apathy, mutism)

Herpes Simplex Virus Infection

Pathogenesis. Herpes simplex virus type 1 (HSV-1) is usually transmitted in childhood through lesions of the oral mucosa (gingival stomatitis, pharyngitis). The virus travels centripetally by way of nerve processes toward the sensory ganglia (e.g., the trigeminal ganglion), where it remains dormant for a variable period of time until reactivated by a trigger such as ultraviolet radiation, dental procedures, immunosuppression, or a febrile illness. It then travels centrifugally, again over nerve processes, back to the periphery, producing blisterlike vesicles (herpes labialis). HSV-1 also causes eye infection (keratoconjunctivitis), as well as (meningo)encephalitis when it spreads via CN I and leptomeningeal fibers of CN V. There is no association between herpes labialis and HSV-1 encephalitis. Herpes simplex virus type 2 (HSV-2) reaches the lumbosacral ganglia by axonal transport from a site of (asymptomatic) urogenital infection. Its reactivation causes genital herpes. In adults, HSV-2 infection can cause (aseptic) meningitis and, occasionally, polyradiculitis or myelitis. HSV-2 virus can be transmitted to the newborn during the birth process, causing encephalitis. HSV-1 encephalitis is very rare in neonates, and HSV-2 encephalitis is very rare in adults.

Symptoms and signs. Herpes simplex encephalitis (HSE) in adults begins with local inflammation of the caudal and medial parts of the frontal and temporal lobes. Uncharacteristic prodromal signs such as fever, headache, nausea, anorexia, and lethargy last a few days at the most. Focal symptoms including olfactory and gustatory hallucinations, aphasia, and behavioral disturbances (confusion, psychosis) then appear, along with focal or complex partial seizures with secondary generalization. There is usually repeated seizure activity, but status epilepticus is rare. Intracranial hypertension causes impairment of consciousness or coma within a few hours. In neonates, the inflammation spreads throughout the CNS.

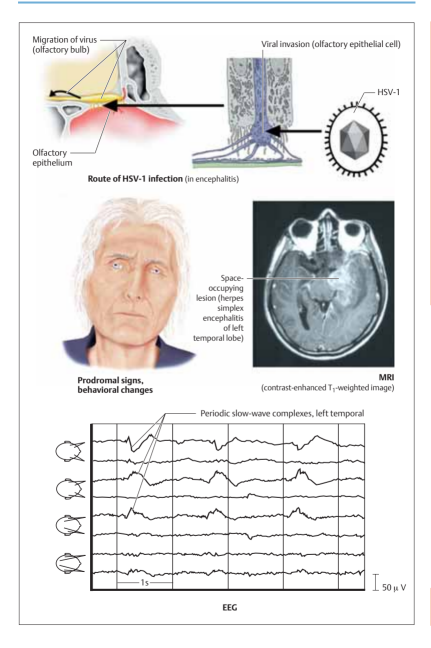
The diagnosis of HSE can be difficult, especially at first. The *clinical findings* include neck stiffness, hemiparesis, and mental disturbances. *CSF examination* reveals a lymphomonocytic pleocytosis (granulocytes may predominate initially) with an elevated protein concentration; low glucose and high lactate concentrations are only rarely found. Xanthochromia and erythrocytes may be present (hemorrhagic necrotizing encephalitis). In the first 3 weeks, the virus can almost always be detected in the CSF by polymerase chain reaction: brain biopsy is only rarely needed for identification of the viral pathogen. Lumbar puncture carries a risk if intracranial hypertension is present (p. 162). EEG reveals periodic high-voltage sharp waves and 2-3 Hz slow wave complexes as a focal or diffuse finding in one or both temporal lobes. In the acute stage of HSE, CT is normal or reveals only mild temporobasal hypointensity without contrast enhancement. Hemorrhage may appear as a hyperdense area. Sharply defined areas of hypodensity appear on CT only in the later stages of HSE. T2-weighted MRI, however, already reveals inflammatory lesions in early HSE. Thus, MRI is used for early diagnosis, CT for the monitoring of encephalitic foci and cerebral edema over the course of the disease.

Meningitis. The clinical manifestations are those of aseptic meningitis (p. 234).

Myelitis. Low back pain, fever, sensory deficit with spinal level, flaccid or spastic paraparesis, bladder and bowel dysfunction. These manifestations usually regress within 2 weeks.

Radiculitis. Inflammation of the lumbosacral nerve roots produces a sensory deficit and bladder and bowel dysfunction.

Virustatic agents. HSV infection of the CNS is treated with acyclovir 10 mg/kg (i.v.) q8h for 14–21 days. Particularly in HSE, it is important to begin treatment as soon as possible.



Varicella-Zoster Virus Infection

Pathogenesis. In children, primary infection with varicella-zoster virus (VZV) usually causes chickenpox (varicella). The portals of entry for infection by droplets or mucus are the conjunctiva, oropharynx, and upper respiratory tract. The virions replicate locally, then enter cells of the reticulohistiocytic system by hematogenous and lymphatic spread (primary viremia). There, they replicate again and disseminate (secondary viremia). VZV infection is followed by immunity. VZV travels by centripetal axonal transport through the sensory nerve fibers of the skin and mucous membranes to the spinal and cranial ganglia and may remain latent there for years (ganglionic latency phase). The thoracic and trigeminal nerve ganglia are most commonly affected, but those of CN VII, IX, and X can also be involved. Spontaneous viral reactivation in the ganglia (ganglionitis) is most common in the elderly, diabetics, and immunocompromised persons (HIV, lymphoma, radiotherapy, chemotherapy, etc.). The reactivated virus travels over the axons centrifugally to the dermatome corresponding to its ganglion of origin, producing the typical dermatomal rash of herpes zoster. It may also spread to the CNS via the spinal dorsal roots (radiculitis), causing herpes zoster myelitis or meningoencephalitis. VZV attacks cerebral blood vessels by way of axonal transport from the trigeminal ganglion. Postherpetic neuralgia is thought to be due to disordered nociceptive processing in both peripheral and central structures.

Symptoms and signs. Chickenpox: After an incubation period of 14-21 days, crops of itchy efflorescent lesions appear, which progress through the sequence macule, papule, vesicle, scab within a few hours. The scabs detach in 1-2 weeks. Immunocompromised patients can develop severe hemorrhagic myelitis, pneumonia, encephalitis, or hepatitis. Acute cerebellitis in children causes appendicular, postural, and gait ataxia, less commonly dysarthria and nystagmus. CSF examination reveals mild pleocytosis and elevation of the protein concentration, or is normal, and the MRI is usually normal. VZV cerebellitis resolves slowly in most cases.

Herpes zoster begins with general symptoms (lethargy, fever) followed by pain, itching, burn-

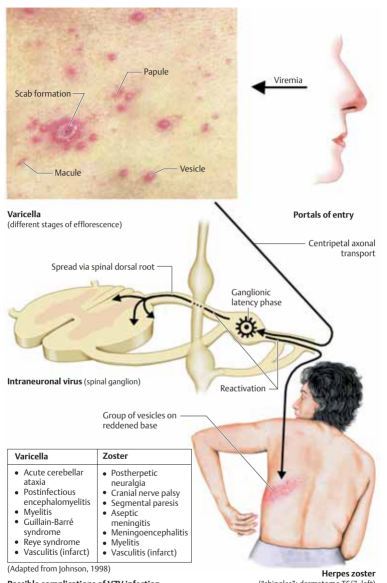
ing or tingling in the affected dermatome(s), which are most commonly thoracic or craniocervical (special forms: *herpes zoster ophthalmicus, oticus,* and *occipitocollaris*). Within a few days, groups of distended vesicles containing clear fluid appear on an erythematous base within the affected dermatome. The contents of the vesicles become turbid and yellowish in 2–3 days. The rash dries, becomes encrusted, and heals in another 5–10 days. The pain and dysesthesia of herpes zoster generally last no longer than 4 weeks. They may also occur without a rash (*herpes zoster sine herpete*).

Complications. Elderly and immunocompromised persons are at increased risk for complications. Pain that persists more than 4 weeks after the cutaneous manifestations have healed is called postherpetic neuralgia and is most common in the cranial and thoracic dermatomes. Cranial nerve involvement may cause unilateral or bilateral ocular complications (ophthalmoplegia, keratoconjunctivitis, visual impairment) or Ramsay-Hunt syndrome (facial palsy, hearing loss, tinnitus, vertigo). Other cranial nerves (IX, X, XII) are rarely affected. Further complications include Guillain-Barré syndrome, myelitis, segmental muscular paresis/ atrophy, myositis, meningitis, ventriculitis, encephalitis, autonomic disturbances (anhidrosis, complex regional pain syndrome), generalized herpes zoster, and vasculitis (ICA and its branches, basilar artery). The viral pathogen is detected in CSF with the polymerase chain reaction.

Virustatic therapy. Acyclovir: 5 mg/kg i. v. q8h or 800 mg p.o. 5 times daily; brivudine: 125 mg p.o. 4 times daily; famcyclovir: 250 mg p.o. 3 times dialy; or valacyclovir: 1 g p.o. 3 times daily. Treatment is continued for 5–7 days. These agents are only effective during the viral replication phase. Intrathecal administration of methylprednisolone is effective in postherpetic neuralgia.

Central Nervous System

CNS Infections



Possible complications of VZV infection

("shingles"; dermatome T6/7, left)

Human Immunodeficiency Virus (HIV) Infection

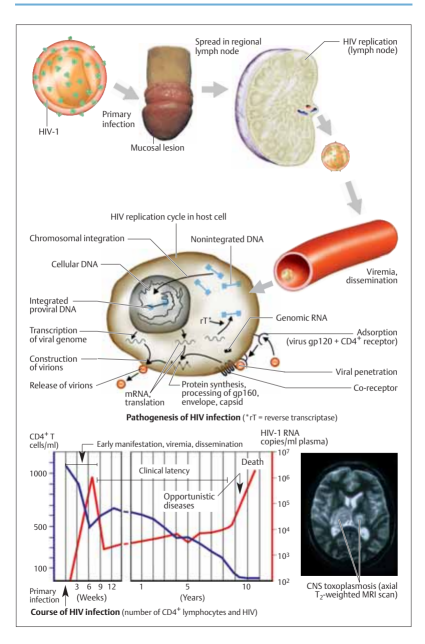
Pathogenesis. HIV type 1 (HIV-1) is found worldwide, HIV-2 mainly in western Africa and only rarely in Europe, America, and India. HIV is transmitted by sexual contact, by exposure to contaminated blood or blood products, or from mother to neonate (vertical transmission). It is not transmitted through nonsexual contact during normal daily activities, by contaminated food or water, or by insect bites. In industrialized countries, the mean incubation period for HIV is 9-12 years, and the mean survival time after the onset of acquired immunodeficiency syndrome (AIDS) is 1-3 years. In primary infection (transmitted through mucosal lesions, etc.), the free or cell-bound organisms enter primary target cells in the hematopoietic system (T cells, B cells, macrophages, dendritic cells), CNS (macrophages, microglia, astrocytes, neurons), skin (fibroblasts), or gastrointestinal tract (goblet cells). After replicating in the primary target cells, the virions spread to regional lymph nodes, CD4+ T cells, and macrophages, where they replicate rapidly, leading to a marked viremia with dissemination of HIV to other target cells throughout the body. About 7-14 days after this viremic phase, the immune system gains partial control over viral replication, and seroconversion occurs. The subsequent period of clinical latency is characterized by a steady rate of viral replication, elimination of HIV by the immune system, and the absence of major clinical manifestations for several years. Eventually, the immune system fails to keep up with the replicating virus, various immune functions become impaired, and the CD4+ T-lymphocyte count declines sharply. The rising viral load correlates with the progression of HIV infection to AIDS. In the nervous system, HIV initially appears in the CSF, but is later found mainly in macrophages and microglia.

Symptoms and signs. Neurological manifestations can occur at any stage of HIV infection, but usually appear only in the late stages of AIDS. One-half to two-thirds of all HIV-positive individuals develop neurological disturbances as a primary or secondary complication of HIV infection or of a concomitant disease.

Primary HIV infection. Early manifestations at the time of seroconversion are rare; these in-

clude acute reversible encephalitis or aseptic meningitis (p. 234), cranial nerve deficits (especially facial nerve palsy), radiculitis, or myelitis. Neurological signs are usually late manifestations of HIV infection. HIV encephalopathy progresses over several months and is characterized by lethargy, headache, increasing social withdrawal, insomnia, forgetfulness, lack of concentration, and apathy. Advanced AIDS is accompanied by bradyphrenia, impaired ocular pursuit, dysarthrophonia, incoordination, myoclonus, rigidity, and postural tremor. Incontinence and central paresis develop in the final stages of the disease. CT of the brain reveals generalized atrophy, and MRI reveals multifocal or diffuse white-matter lesions. The CSF examination may be normal or reveal a low-grade pleocytosis and an elevated protein concentration. EEG reveals increased slow-wave activity. Other neurological manifestations include HIV myelopathy (vacuolar myelopathy), distal symmetrical polyradiculoneuropathies, mononeuritis multiplex, and polymyositis.

Secondary complications of HIV infection include opportunistic CNS infections (toxoplasmosis, cryptococcal meningitis, aspergillosis, progressive multifocal leukoencephalopathy, cytomegalovirus encephalitis, herpes simplex encephalitis, herpes zoster, tuberculosis, syphilis), *tumors* (primary CNS lymphoma), *stroke* (infarction, hemorrhage), and *metabolic disturbances* (iatrogenic or secondary to vitamin deficiency). Virustatic treatment. Antiretroviral combination therapy (HAART: highly active antiretroviral therapy).



Poliomyelitis

Pathogenesis. There are three types of poliovirus: type 1 (ca. 85% of all infections), type 2 and type 3. Like other enteroviruses (e.g., coxsackievirus, echovirus, and hepatitis-A virus), they are transmitted via the fecal-oral and oraloral routes, and poor sanitary conditions favor their spread. Having entered the body, the virions infiltrate epithelial cells, where they replicate, and then spread to the lymphatic tissues of the nasopharynx (tonsils) and intestinal wall (Peyer's patches). A second replication phase (6-8 days) is followed by hematogenous dissemination (viremia), with nonspecific symptoms. Polioviruses reach the CNS via the bloodstream and can produce signs of poliomyelitis 10-14 days after infection. The virus is present in the saliva for 3-4 days and in the feces for 3-4 weeks. The infected individual becomes immune only to the specific type of poliovirus that caused the infection. The viral pathogen can be detected in throat smears and feces by serology or by the polymerase chain reaction.

Symptoms and signs. 90–95% of all poliovirus infections remain asymptomatic (occult immunization). Roughly 5–10% of infected persons develop abortive poliomyelitis, while only 1–2% go on to develop major spinal, bulbar, or encephalitic disease.

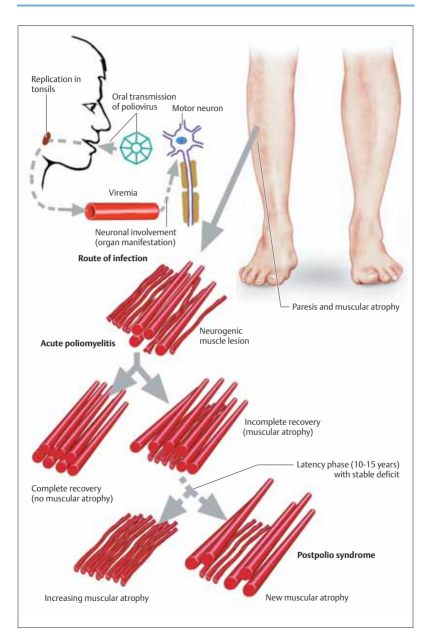
Minor poliomyelitis (abortive type) has nonspecific manifestations including fever, headache, sore throat, limb pain, lethargy, and gastrointestinal disturbances (nausea, anorexia, diarrhea, constipation), which resolve in 4 days at most, without CNS involvement.

Major poliomyelitis (preparalytic and paralytic types). Either immediately or after a latency period of 2–3 days, fever rises and organ manifestations appear, with a meningitic syndrome (preparalytic stage) that may be followed by paralysis in the later course of the disease (paralytic stage). The meningitis of the preparalytic stage exhibits typical features of aseptic meningitis as well as marked generalized weakness and apathy. It resolves in about one-half of cases; in the other half, increasing myalgia and stiffness herald the onset of the paralytic stage. The spinal form (most common) causes flaccid paresis (usually with asymmetrical proximal weakness) and areflexia, mainly in the lower limbs. The paresis may worsen over 3-5 days; its severity is highly variable. Some patients develop paresthesiae without sensory loss or autonomic dysfunction (urinary retention, hypohidrosis. constipation). Muscle atrophy develops within a week of the onset of paralysis. Bulbar poliomyelitis develops in some 10% of patients (in isolated form, or concurrently with spinal poliomyelitis), involving CN VII, IX, and X to produce dysphagia and dysphonia. Involvement of the brain stem reticular formation causes hemodynamic fluctuations, respiratory insufficiency or paralysis, and gastric atony. The encephalitic form is very rare; it may be accompanied by autonomic dysfunction (p. 222).

Postpolio syndrome. Newly arising manifestations in a patient who recovered from poliomyelitis at least 10 years earlier with stable neurological deficits in the intervening time. Postpolio syndrome is characterized by general symptoms (abnormal fatigability, intolerance to cold, cyanosis of the affected limbs, etc.), arthralgias, and increasing neuromuscular deficits (exacerbation of earlier weakness, weakness of previously unaffected muscles, new atrophy), sometimes accompanied by dysphagia, respiratory insufficiency, and sleep apnea.

Prevention. Subcutaneous immunization with inactivated polioviruses (e. g., Salk vaccine), followed by a first booster in 6–8 weeks and a second booster in 8–12 months.

CNS Infections



Progressive Multifocal Leukoencephalopathy (PML)

Pathogenesis. The causative organism, JC virus, is a ubiquitous papovavirus that usually stays dormant within the body. It is reactivated in persons with impaired cellular immunity and spreads through the bloodstream to the CNS, where it induces multiple white-matter lesions. Symptoms and signs. PML appears as a complication of cancer (chronic lymphatic leukemia, Hodgkin lymphoma), tuberculosis, sarcoidosis, immune suppression, and AIDS, producing variable symptoms and signs. The major manifestations in patients without AIDS are visual disturbances (visual field defects, cortical blindness), hemiparesis, and neuropsychological disturbances (impairment of memory and cognitive functions, dysphasia, behavioral abnormalities). The major manifestations of PML as a complication of AIDS are (from most to least frequent): central paresis, cognitive impairment, visual disturbances, gait impairment, ataxia, dysarthria, dysphasia, and headache. PML usually progresses rapidly, causing death in 4-6 months. The definitive diagnosis is by histological examination of brain tissue obtained by biopsy or necropsy. CT reveals asymmetrically distributed, hypodense white-matter lesions without mass effect or contrast enhancement; these lesions are hyperintense on T2-weighted MRI, which also demonstrates involvement of the subcortical white matter ("U fibers"). The CSF findings are usually normal, but oligoclonal bands may be found in AIDS patients.

Virustatic therapy. There is as yet no validated treatment regimen.

Cytomegalovirus (CMV) Infection

Pathogenesis. CMV, a member of the herpesvirus family, is transmitted through respiratory droplets, sexual intercourse, and contact with contaminated blood, blood products, or transplanted organs. It is widely distributed throughout the world, with a regional and agedependent prevalence of up to 100%. CMV virions are thought to replicate initially in oropharyngeal epithelial cells (salivary glands) and then disseminate to the organs of the body, including the nervous system, through the bloodstream. The virus remains dormant in monocytes and lymphocytes as long as the immune system keeps it in check. Reactivation of the virus is almost always asymptomatic in healthy individuals, but severe generalized disease can develop in persons with immune compromise due to AIDS, organ transplantation, immunosuppressant drugs, or a primary malignancy.

Symptoms and signs. The primary infection is usually clinically silent. Intrauterine fetal infection leads to generalized fetopathies in fewer than 5% of neonates. In immunocompromised patients, particularly those with AIDS, (reactivated) CMV infection presents a variable combination of manifestations, including retinitis (partial or total loss of vision), pneumonia, and enteritis (colitis, esophagitis, proctitis). The neurological manifestations of CMV infection are manifold. PNS involvement is reflected as Guillain-Barré syndrome or lumbosacral polyradiculopathy (subacute paraparesis with or without back pain or radicular pain). CNS involvement produces encephalitis, meningitis, ventriculitis (inflammatory changes in the ependyma) and/or myelitis. Symptoms and signs may be absent, minor, or progressively severe, as in HIV-related encephalopathy. CMV vasculitis may lead to ischemic stroke. The diagnosis usually cannot be made from the clinical findings alone (except in the case of CMV retinitis). MRI reveals periventricular contrast enhancement in CMV vasculitis; other MRI and CT findings are nonspecific. There may be CSF pleocytosis with an elevated protein concentration. The diagnosis can be established by culture or identification by polymerase chain reaction of CMV in tissue, CSF, or urine, or by serological detection of CMV-specific antibodies.

Virustatic therapy. Gancyclovir, foscarnet, or cidofovir are given for initial treatment and secondary prophylaxis.

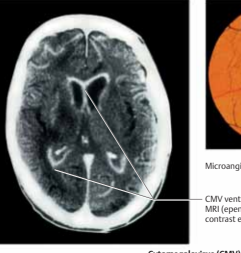


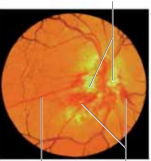
Foci of demyelination seen on MRI (no mass effect or contrast enhancement)

Dysarthria, dysphasia, cognitive impairment, behavioral changes

Progressive multifocal leukoencephalopathy

Cotton-wool spots near optic disk





Microangiopathy Hemorrhage CMV retinitis

 CMV ventriculitis on MRI (ependymal contrast enhancement)

Cytomegalovirus (CMV) infection

Rabies

Pathogenesis. Rabies virus is a rhabdovirus that is mainly transmitted by the bite of a rabid animal. The reservoirs of infection are wild animals in Europe and America (foxes, wild boar, deer, martens, raccoons, badgers, bats; sylvatic rabies) and dogs in Asia (urban rabies). The virus replicates in muscles cells near the site of entry and then spreads via muscle spindles and motor end plates to the peripheral nerves, as far as the spinal ganglia and spinal motor neurons, where secondary replication takes place. It subsequently spreads to the CNS and other organs (salivary glands, cornea, kidneys, lungs) by way of the fiber pathways of the autonomic nervous system. The limbic system (p. 144) is usually also involved. The mean incubation time is 2-3 months (range: 1 week to 1 year). Proof that the biting animal was rabid is essential for diagnosis, as rabies is otherwise very difficult to diagnose until its late clinical manifestations appear. The virus can be isolated from the patient's sputum, urine or CSF in the first week after infection.

Symptoms and signs. The course of rabies can be divided into three stages. The *prodromal stage* (2–4 days) is characterized by paresthesia, hyperesthesia, and pain at the site of the bite and the entire ipsilateral side of the body. The patient suffers from nausea, malaise, fever, and headache and, within a few days, also from anxiety, irritability, insomnia, motor hyperactivity, and depression.

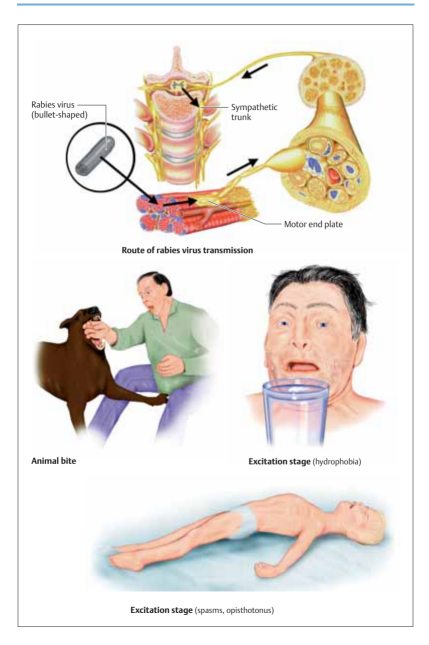
Hyperexcitability stage. In the ensuing days, the patient typically develops increasing restlessness, incoherent speech, and painful spasms of the limbs and muscles of deglutition, reflecting involvement of the midbrain tegmentum. Hydrophobia, as this stage of the disease is called, is characterized by painful laryngospasms, respiratory muscle spasms, and opisthotonus, with tonic-clonic spasms throughout the body that are initially triggered by attempts to drink but later even by the mere sight of water, unexpected noises, breezes, or bright light. There may be alternating periods of extreme agitation (screaming, spitting, and/or scratching fits) and relative calm. The patient dies within a few days if untreated, or else progresses to the next stage after a brief clinical improvement.

Paralytic stage (paralytic rabies). The patient's mood and hydrophobic manifestations improve, but spinal involvement produces an ascending flaccid paralysis with myalgia and fasciculations. Weakness may appear in all limbs at once, or else in an initially asymmetrical pattern, beginning in the bitten limb and then spreading. In some cases, the clinical picture is dominated by cranial nerve palsies (oculomotor disturbances, dysphagia, drooling, dysarthrophonia) and autonomic dysfunction (cardiac arrhythmia, pulmonary edema, diabetes insipidus, hyperhidrosis).

Rabies prophylaxis. *Preexposure prophylaxis:* Vaccination of persons at risk (veterinarians, laboratory personnel, travelers to endemic areas).

Local wound treatment: Thorough washing of the bite wound with soap and water.

Postexposure prophylaxis: Vaccination and rabies immunoglobulin.



Opportunistic Fungal Infections

CNS mycosis is sometimes found in otherwise healthy persons but mainly occurs as a component of an opportunistic systemic mycosis in persons with immune compromise due to AIDS, organ transplantation, severe burns, malignant diseases, diabetes mellitus, connective tissue diseases, chemotherapy, or chronic corticosteroid therapy. Certain types of mycosis (blastomycosis, coccidioidmycosis, histoplasmosis) are endemic to certain regions of the world (North America, South America, Africa).

Cryptococcus neoformans (Cryptococcosis)

Cryptococcus, a yeastlike fungus with a polysaccharide capsule, is a common cause of CNS mycosis. It is mainly transmitted by inhalation of dust contaminated with the feces of pet birds and pigeons. Local pulmonary infection is followed by hematogenous spread to the CNS. In the presence of a competent immune system (particularly cell-mediated immunity), the pulmonary infection usually remains asymptomatic and self-limited. Immune-compromised persons, however, may develop meningoencephalitis with or without prior signs of pulmonary cryptococcosis. Its manifestations are heterogeneous and usually progressive. Signs of subacute or chronic meningitis are accompanied by cranial nerve deficits (III, IV, VI), encephalitic syndrome, and/or signs of intracranial hypertension. Diagnosis: MRI reveals granulomatous cystic lesions with surrounding edema. Lung infiltrates may be seen. The nonspecific CSF changes include a variable (usually mild) lymphomonocytic pleocytosis as well as elevated protein, low glucose, and elevated lactate concentrations. An india ink histological preparation reveals the pathogen with a surrounding halo (carbon particles cannot penetrate its polysaccharide capsule). Identification of pathogen: demonstration of antigen in CSF and serum; tests for anticryptococcal antibody yield variable results. Treatment: initially, amphotericin B + flucytosine; subsequently, fluconazole or (if fluconazole is not tolerated) itraconazole.

Candida (Candidiasis)

Candida albicans is a constituent of the normal body flora. In persons with impaired cell-medi-

ated immunity, Candida can infect the oropharynx (thrush) and then spread to the upper respiratory tract, esophagus, and intestine. CNS infection comes about by hematogenous spread (candida sepsis), resulting in meningitis or meningoencephalitis. Ocular changes: Candida endophthalmitis. Diagnosis: Candida abscesses can be seen on CT or MRI. The CSF changes included pleocytosis (several hundred cells/ul) and elevated concentrations of protein and lactate. Pathogen identification: Microscopy, culture, or detection of specific antigens or antibodies. Local treatment: Amphotericin B or fluconazole. Systemic tratment: Amphotericin B + flucytosine.

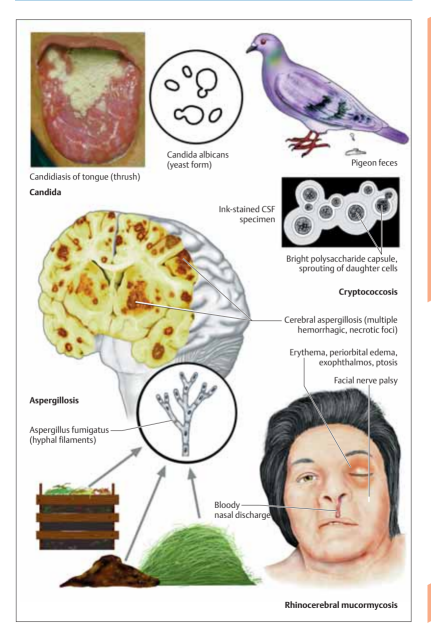
Aspergillus (Aspergillosis)

The mold Aspergillus fumigatus is commonly found in cellulose-containing materials such as silage grain, wood, paper, potting soil, and foliage. Inhaled spores produce local inflammation in the airways, sinuses, and lungs. Organisms reach the CNS by hematogenous spread or by direct extension (e.g., from osteomyelitis of the skull base, otitis, or mastoiditis), causing encephalitis, dural granulomas, or multiple abscesses. Diagnosis: CT and MRI reveal multiple, sometimes hemorrhagic lesions. The CSF findings include granulocytic pleocytosis and markedly elevated protein, decreased glucose, and elevated lactate concentration. Pathogen identification: Culture; if negative, then lung or brain biopsy. Treatment: Amphotericin B + flucytosine or itraconazole.

Mucor, Absidia, Rhizopus (Mucormycosis)

Inhaled spores of these molds enter the nasopharynx, bronchi, and lungs, where they mainly infect blood vessels. *Rhinocerebral mucormycosis* is a rare complication of diabetic ketoacidosis, lymphoproliferative disorders, and drug abuse; infection spreads from the paranasal sinuses via blood vessels to the retro-orbital tissues (causing retro-orbital edema, exophthalmos, and ophthalmoplegia) and to the brain (causing infarction with secondary hemorrhage). *Diagnosis*: CT, MRI; associated findings on ENT examination. *Pathogen identification*: Biopsy, smears. *Treatment*: Surgical excision of infected tissue if possible; amphotericin B.

CNS Infections



Protozoan and Helminthic Infections

Toxoplasma gondii (Toxoplasmosis)

This protozoan goes through three stages of development. Tachyzoites (endozoites; acute stage) are crescent-shaped, rapidly replicating forms that circulate in the bloodstream and are spread from one individual to another through contaminated blood or blood products. These develop into bradyzoites (cystozoites; latent stage), which aggregate to form tissue cysts (e.g., in muscle) containing several thousand organisms each. Oocysts are found only in the intestinal mucosa of the definitive host (domestic cat). Infectious sporozoites (sporulated oocysts) appear 2-4 days after the oocysts are eliminated in cat feces. Reuptake of the organism by the definitive host, or infection of an intermediate host (human, pig, sheep), occurs by ingestion of sporozoites from contaminated feces, or by consumption of raw meat containing tissue cysts. In the intermediate host, the sporozoites develop into tachyzoites, which then become bradyzoites and tissue cysts. Placental transmission (congenital toxoplasmosis \Rightarrow hydrocephalus, intracellular calcium deposits, chorioretinitis) occurs only if the mother is initially infected during pregnancy. In immunocompetent persons, acute toxoplasmosis is usually asymptomatic, and only occasionally causes symptoms such as lymphadenopathy, fatigue, low-grade fever, arthralgia, and headache. IgG antibodies can be detected in latent toxoplasmosis (bradyzoite stage). In immunodeficient persons (p. 240), however, latent toxoplasmosis usually becomes symptomatic on reactivation. The central nervous system is most commonly affected (mainly encephalitis; myelitis is rare); other organs that may be affected include the eyes (chorioretinitis, iridocyclitis), heart, liver. spleen, PNS (neuritis) and muscles (myositis).

Diagnosis: EEG (slowing, focal signs), CT/MRI (solitary or multiple ring-enhancing abscesses), CSF (lymphomonocytic pleocytosis, mildly elevated protein concentration). Treatment: pyrimethamine/sulfadiazine or clindamycin/ folinic acid.

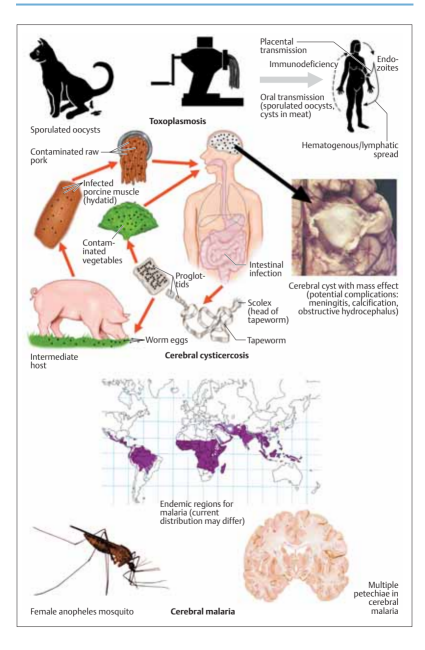
Taenia solium (Neurocysticercosis)

Ingestion of the tapeworm *Taenia solium* in raw or undercooked pork leads to a usually asymp-

tomatic infection of the human gut. Tapeworm segments that contain eggs (proglottids) are eliminated in the feces of pigs (the intermediate host) or humans with intestinal infection and then reingested by humans (or pigs) under poor hygienic conditions. The oval-shaped larvae pass through the intestinal wall and travel to multiple organs (including the eyes, skin, muscles, lung, and heart) by hematogenous, lymphatic, or direct spread. The CNS is often involved, though manifestations such as epileptic seizures, intracranial hypertension, behavioral changes (dementia, disorientation), hemiparesis, aphasia, and ataxia are uncommon. Spinal cysts are rare. Diagnosis: CT (solitary or multiple hypodense cysts with or without contrast enhancement. calcification and/or hvdrocephalus), MRI (demonstration of cysts and surrounding edema), CSF examination (low-grade lymphocytic pleocytosis, occasional eosinophilia). Treatment: Praziguantel or albendazole; neurosurgical excision of intraventricular cysts; ventricular shunting in patients with hydrocephalus.

Plasmodium falciparum (Cerebral Malaria)

This protozoan is most commonly transmitted by the bite of the female anopheles mosquito. Primary asexual reproduction of the organisms takes place in the hepatic parenchyma (*preerythrocytic schizogony*). The organisms then invade red blood cells and develop further inside them (*intraerythrocytic development*). The repeated liberation of merozoites causes recurrent episodes of fever. *P. falciparum* preferentially colonizes the capillaries of the brain, heart, liver, and kidneys. *Pathogen identification:* Blood culture. *Treatment:* See current topical literature for recommendations.



Transmissible Spongiform Encephalopathies

The transmissible spongiform encephalopathies (TSEs) are characterized by spongiform histological changes in the brain (vacuoles in neurons and neuropil), transmissibility to humans by way of infected tissue or contaminated surgical instruments, and, in some cases, a genetic determination. TSEs are transmitted by nucleic acidfree proteinaceous particles called prions and are associated with mutations in prion protein (PrP); they are therefore referred to as *prion diseases*.

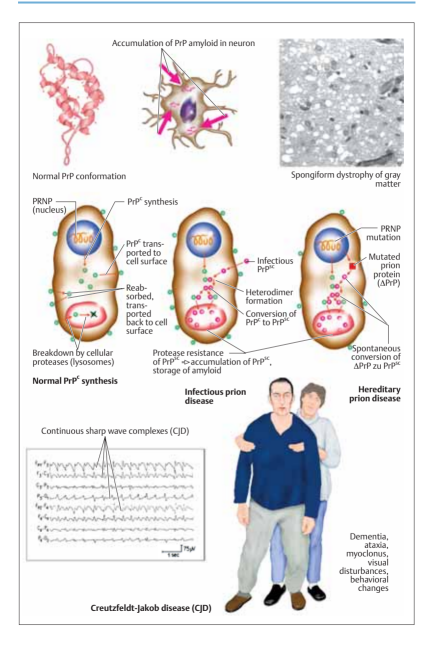
Normal cellular prion protein (PrPc) is synthesized intracellularly, transported to the cell membrane, and returned to the cell interior by endocytosis. Part of the PrPc is then broken down by proteases, and another fraction is transported back to the cell surface. The physiological function of PrPc is still unknown. It is found in all mammalian species and is especially abundant in neurons. PRNP, the gene responsible for the expression of PrP^c in man, is found on the short arm of chromosome 20. PRNP mutations yield the mutated form of PrP (Δ PrP) that causes the genetic spongiform encephalopathies. Another mutated form of PrP (PrPsc) causes the infectious spongiform encephalopathies. PrPsc induces the conversion of PrPc to PrPsc in the following manner: PrPsc enters the cell and binds with PrPc to yield a heterodimer. The resulting conformational change in the PrPc molecule (a-helical structure) and its interaction with a still unidentified cellular protein (protein X) transform it into PrPsc (β-sheet structure). Protein X is thought to supply the energy needed for protein folding, or at least to lower the activation energy for it. PrPsc cannot be formed in cells lacking PrPc. Mutated PrPsc presumably reaches the CNS by axonal transport or in lymphatic cells; these forms of transport have been demonstrated in forms of spongiform encephalopathies that affect domestic animals, e.g., scrapie (in sheep) and bovine spongiform encephalopathy (BSE). ΔPrP and PrPsc cannot be broken down intracellularly and therefore accumulate within the cells. Partial proteolysis of these proteins yields a protease-resistant molecule (PrP 27-30) that polymerizes to form amyloid, which, in turn, induces further neuropathological changes. PrP and amyloid have been found in certain myopathies (such as inclusion body myositis, p. 344); others involve an accumulation of PrP (PrP overexpression myopathy).

Creutzfeldt–Jakob Disease (CJD)

CID is a very rare disease, arising in ca. 1 person per 106 per year. It usually affects older adults (peak incidence around age 60). 85-90% of cases are sporadic (due to a spontaneous gene mutation or conformational change of PrPc to PrPsc); 5-15% are familial (usually autosomal dominant); and very rare cases are iatrogenic (transmitted by contaminated neurosurgical instruments or implants, growth hormone, and dural and corneal grafts). It usually progresses rapidly to death within 4-12 months of onset, though the survival time in individual cases varies from a few weeks to several years. Early manifestations are not typically seen, but may include fatigability, vertigo, cognitive impairment, anxiety, insomnia, hallucinations, increasing apathy, and depression. The principal finding is a rapidly progressive dementia associated with myoclonus, increased startle response, motor disturbances (rigidity, muscle atrophy, fasciculations, cerebellar ataxia), and visual disturbances. Late manifestations include akinetic mutism, severe myoclonus, epileptic seizures, and autonomic dysfunction. A new variant of CJD has recently arisen in the United Kingdom; unlike the typical form, it tends to affect younger patients, produces mainly behavioral changes in its early stages, and is associated with longer survival (though it, too, is fatal). It is thought to be caused by the consumption of beef from cattle infected with BSE. Diagnosis: EEG (1 Hz periodic biphasic or triphasic sharp-wave complexes), CT (cortical atrophy), T2/proton-weighted MRI (bilateral hyperintensity in basal ganglia in ca. 80%), CSF examination (elevation of neuron-specific enolase, S100^β or tau protein concentration; presence of protein 14-3-3).

Gerstmann-Sträussler-Scheinker Disease (GSS) and Fatal Familial Insomnia (FFI)

See pp. 114 and 280.



Symptoms and Signs

The clinical manifestations of a brain tumor may range from a virtually asymptomatic state to a constellation of symptoms and signs that is specific for a particular type and location of lesion. The only way to rule out a brain tumor for certain is by neuroimaging (CT or MRI).

Nonspecific Manifestations

Tumors whose manifestations are mainly nonspecific include astrocytoma, oligodendroglioma, cerebral metastasis, ependymoma, meningioma, neoplastic meningitis, and primary CNS lymphoma.

Behavioral changes. Patients may complain of easy fatigability or exhaustion, while their relatives or co-workers may notice lack of concentration, forgetfulness, loss of initiative, cognitive impairment, indifference, negligent task performance, indecisiveness, slovenliness, and general slowing of movement. Such manifestations are often mistaken for signs of depression or stress. Apathy, obtundation, and somnolence worsen as the disease progresses. There may also be increasing confusion, disorientation, and dementia.

Headache. More than half of patients with brain tumors suffer from headache, and many headache patients fear that they might have a brain tumor. If headache is the sole symptom, the neurological examination is normal, and the headache can be securely classified as belonging to one of the primary types (p. 182 ff), then a brain tumor is very unlikely. Neuroimaging is indicated in patients with longstanding headache who report a change in their symptoms. The clinical features of headache do not differentiate benign from malignant tumors.

Nausea, vertigo, and malaise are frequent, though often vague, complaints. The patient feels unsteady or simply "different." Vomiting (sometimes on an empty stomach) is less common and not necessarily accompanied by nausea; there may be spontaneous, projectile vomiting.

Epileptic seizures. Focal or generalized seizures arising in adulthood should prompt evaluation for a possible brain tumor.

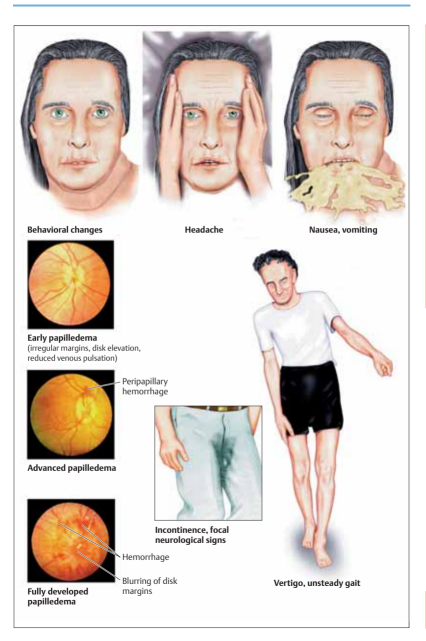
Focal neurological signs usually become prominent only in advanced stages of the disease but may be present earlier in milder form. Hemiparesis, aphasia, apraxia, ataxia, cranial nerve palsies, or incontinence may occur depending on the type and location of the tumor.

Intracranial hypertension (elevated ICP) (p. 158) may arise without marked focal neurological dysfunction because of a medulloblastoma, ependymoma of the fourth ventricle, cerebellar hemangioblastoma, colloid cyst of the 3rd ventricle, craniopharyngioma, or glioblastoma (e.g. of the frontal lobe or corpus callosum). Cervical tumors very rarely cause intracranial hypertension. Papilledema, if present, is not necessarily due to a brain tumor, nor does its absence rule one out. Papilledema does not impair vision in its acute phase.

Specific Manifestations

Some tumors produce symptoms and signs that are specific for their histological type, location, or both. These tumors include craniopharyngioma, olfactory groove meningioma, pituitary tumors, cerebellopontine angle tumors, pontine glioma, chondrosarcoma, chordoma, glomus tumors, skull base tumors, and tumors of the foramen magnum. In general, these specific manifestations are typically found when the tumor is relatively small and are gradually overshadowed by nonspecific manifestations (described above) as it grows.

Brain Tumors



Benign Brain Tumors

Astrocytoma (WHO grades I and II)

Astrocytomas arise from blastomatous astrocytes. They are classified as benign (WHO grade I) or semibenign (WHO grade II) according to their histological features (p. 377).

Pilocytic astrocytoma (WHO grade I) is a slowly growing tumor that mainly occurs in children and young adults and usually arises in the cerebellum, optic nerve, optic chiasm, hypothalamus, or pons. It is not uncommonly found in the setting of neurofibromatosis I. There may be a relatively long history of headache, abnormal gait, visual impairment, diabetes insipidus, precocious puberty, or cranial nerve palsies before the tumor is discovered.

Low-grade astrocytoma (WHO grade II). Fibrillary astrocytoma is more common than the gemistocytic and protoplasmic types. These tumors most commonly arise in the frontal and temporal lobes and often undergo malignant transformation to grades III and IV over the course of several years. They may be calcified. They may produce epileptic seizures and behavioral changes.

Oligodendroglioma (WHO grade II) usually appears in the 4th or 5th decade of life. It tends to arise at or near the cortical surface of the frontal and temporal lobes and may extend locally to involve the leptomeninges. Oligodendrogliomas are often partially calcified. Tumors of mixed histology (oligodendrocytoma plus astrocytoma) are called *oligoastrocytomas*.

Pleomorphic xanthoastrocytoma (WHO grade II) is a rare tumor that mainly arises in the temporal lobes of children and young adults and is associated with epileptic seizures in most cases. It can progress to a grade III tumor.

Meningioma (WHO grade I)

Meningiomas are slowly growing, usually benign, dural-based extraaxial tumors that are thought to arise from arachnoid cells. Twelve histological subtypes have been identified. Meningiomas tend to recur if they are not totally resected. They may involve not only the dura mater but also the adjacent bone (manifesting usually as hyperostosis, more rarely as thinning) and may infiltrate or occlude the cerebral venous sinuses. They can occur anywhere in the CNS but are most often found in *supratentorial* (falx, parasagittal region, sphenoid wing, cerebral convexities), *infratentorial* (tentorium, cerebellopontine angle, craniocervical junction), and *spinal* locations. Multiple or intraventricular meningioma is less common. *Extracranial meningiomas* rarely arise in the orbit, skin, or nasal sinuses. Familial meningioma is seen in hereditary disorders such as type II neurofibromatosis.

Choroid Plexus Papilloma (WHO grade I)

This rare tumor most commonly arises in the (left) lateral ventricle in children and in the 4th ventricle in adults. Signs of intracranial hypertension, due to obstruction of CSF flow, are the most common clinical presentation and may arise acutely.

Hemangioblastoma (WHO grade I)

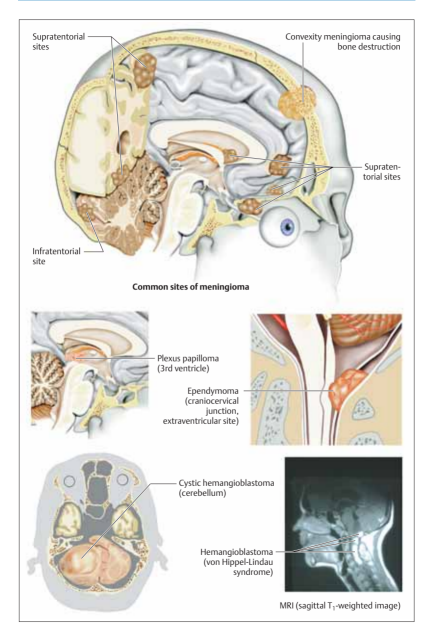
These solid or cystic tumors usually arise in the cerebellum (from the vermis more often than the hemispheres) and produce vertigo, head-ache, truncal ataxia, and gait ataxia. Obstructive hydrocephalus may occur as an early manifestation. 10% of cases are in patients with von Hippel-Lindau disease (p. 294).

Ependymoma (WHO grades I–II)

Ependymomas most commonly arise in children, adolescents, and young adults. They may arise in the ventricular system (usually in the fourth ventricle) or outside it; they may be cystic or calcified. *Subependymoma* of the fourth ventricle (WHO grade 1) may appear in middle age or later. Spinal ependymomas may arise in any portion of the spinal cord.

Central Nervous System

Brain Tumors



257

Tumors in Specific Locations

Supratentorial Region

Colloid cyst of 3rd ventricle. These cysts filled with gelatinous fluid are found in proximity to the interventricular foramen (of Monro). Small colloid cysts may remain asymptomatic, but large ones cause acute or chronic obstructive hydrocephalus (p. 162). Sudden obstruction of the foramen causes acute intracranial hypertension, sometimes with loss of consciousness. Symptomatic colloid cysts can be surgically removed with stereotactic, neuroendoscopic, or open techniques.

Craniopharyngioma (WHO grade I). Adamantinomatous craniopharyngioma is suprasellar tumor of children and adolescents that has both cystic and calcified components. It produces visual field defects, hormonal deficits (growth retardation, thyroid and adrenocortical insufficiency, diabetes insipidus), and hydrocephalus. Large tumors can cause behavioral changes and epileptic seizures. *Papillary* craniopharyngioma is a tumor of adults that usually involves the 3rd ventricle.

Pituitary adenomas (WHO grade I). Adenomas smaller than 10 mm, called microadenomas, are usually hormone-secreting, while those larger than 10 mm, called macroadenomas, are often non-hormone-secreting. In addition to possible hormone secretion, these tumors have intrasellar (hypothyroidism, adrenocortical hormone deficiency, amenorrhea reflecting anterior pituitary insufficiency, and, rarely, diabetes insipidus), suprasellar (chiasmatic lesions, p. 82, hypothalamic compression, hydrocephalus), and parasellar manifestations (headache, deficits of CN III-VI, encirclement of the ICA by tumor, diabetes insipidus), which gradually progress as the tumor enlarges. Hemorrhage or infarction of a pituitary tumor can cause acute pituitary failure (cf. Sheehan's postpartum necrosis of the pituitary gland). Prolactinomas (prolactin-secreting tumors) elevate the serum prolactin concentration above 200 µg/l, in distinction to the less pronounced secondary hyperprolactinemia (usually <200 µg/l) associated with as pregnancy, parasellar tumors, dopamine antagonists (neuroleptics, metoclopramide, reserpine), and epileptic seizures. Prolactinomas can cause secondary amenorrhea, galactorrhea, and hirsutism in women, and headache, impotence, and galactorrhea (rarely) in men. Growth hormone-secreting tumors cause gigantism in adolescents and acromegaly in adults. Headache, impotence, polyneuropathy, diabetes mellitus, organ changes (goiter), and hypertension are additional features. ACTHsecreting tumors cause Cushing disease.

Tumors of the pineal region. The most common tumor of the pineal region is germinoma (WHO grade III), followed by pineocytoma (WHO grade I) and pineoblastoma (WHO grade IV). The clinical manifestations include Parinaud syndrome (p. 358), hydrocephalus, and signs of metastatic dissemination in the subarachnoid space (p. 262).

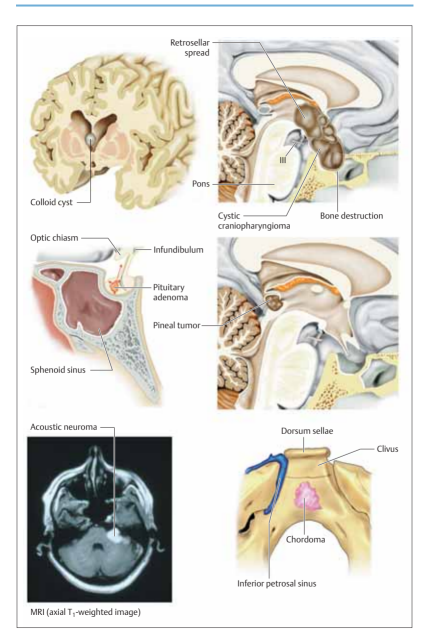
Infratentorial Region

Acoustic neuroma (WHO grade I) is commonly so called, though it is in fact a schwannoma of the vestibular portion of CN VIII. Early manifestations include hearing impairment (rarely sudden hearing loss), tinnitus, and vertigo. Larger tumors cause cranial nerve palsies (V, VII, IX, X), cerebellar ataxia, and sometimes hydrocephalus. Bilateral acoustic neuroma is seen in neurofibromatosis II.

Chordoma arises from the clivus and, as it grows, destroys the surrounding bone tissue and compresses the brain stem, causing cranial nerve palsies (III, V, VI, IX, X, XII), pituitary dysfunction, visual field defects, and headache.

Paragangliomas. This group of tumors includes *pheochromocytoma* (arising from the adrenal medulla), *sympathetic paraganglioma* (arising from neuroendocrine cells of the sympathetic system), and *parasympathetic ganglioma* or *chemodetectoma* (arising from parasympathetically innervated chemoreceptor cells). The lastnamed is a highly vascularized tumor that may grow invasively. It arises from the glomus body.

Brain Tumors



259

Malignant Tumors

Anaplastic Astrocytoma (WHO grade III) and Glioblastoma (WHO grade IV)

These infiltrative, rapidly growing tumors usually arise in adults between the ages of 40 and 65. They usually involve the cerebral hemispheres, but are sometimes found in infratentorial locations (brain stem, cerebellum, spinal cord). They are occasionally multicentric or diffuse (gliomatosis cerebri is extremely rare). Infiltrative growth across the corpus callosum to the opposite side of the head is not uncommon ("butterfly glioma"). These tumors are often several centimeters in diameter by the time of diagnosis. Even relatively small tumors can produce considerable cerebral edema. Metastases outside the CNS (bone, lymph nodes) are rare. CT and MRI reveal ringlike or garlandlike contrast around a hypointense center.

Primary Cerebral Lymphoma (WHO grade IV)

These tumors are usually non-Hodgkin lymphomas of the B-cell type and are only rarely of the T-cell type. They are commonly associated with congenital or acquired immune deficiency (Wiskott-Aldrich syndrome; immune suppression for organ transplantation, AIDS) and can arise in any part of the CNS (80% supratentorial, 20% infratentorial). Headache, cranial nerve palsies, polyradiculoneuropathy, meningismus, and ataxia suggest (primary) leptomeningeal involvement. Ocular manifestations: Infiltration of the uvea and vitreous body (visual disturbances; slit-lamp examination). Moreover, lymphomas may occur as solitary or multiple tumors or may spread diffusely through CNS tissue (periventricular zone, deep white matter). They produce local symptoms and also such general symptoms as psychosis, dementia, and anorexia. CT reveals them as hyperdense lesions surrounded by edema, usually with homogeneous contrast absorption and with little or no mass effect. MRI is more sensitive for lymphoma than CT; it reveals the extent of surrounding edema and is especially useful for the detection of spinal, leptomeningeal, and multilocular involvement. CSF examination reveals malignant cells in the early stages of the disease; the CSF protein concentration is not necessarily elevated. Biopsies and/or CSF serology for diagnosis should be performed before treatment is initiated, because some of the drugs used, particularly corticosteroids, can make the disease more difficult to diagnose.

Anaplastic Oligodendroglioma (WHO grade III)

This rare form of oligodendroglioma responds well to chemotherapy with procarbazine, CCNU, and vincristine (PCV). As histological confirmation of cellular anaplasia (the defining criterion for grade III) can be difficult, the diagnosis must sometimes be based on the clinical and radiological findings. There may be leptomeningeal dissemination or meningeal gliomatosis.

Anaplastic Ependymoma (WHO grade III)

These tumors may have subarachnoid and (rarely) extraneural metastases, e.g., to the liver, lungs and ovaries.

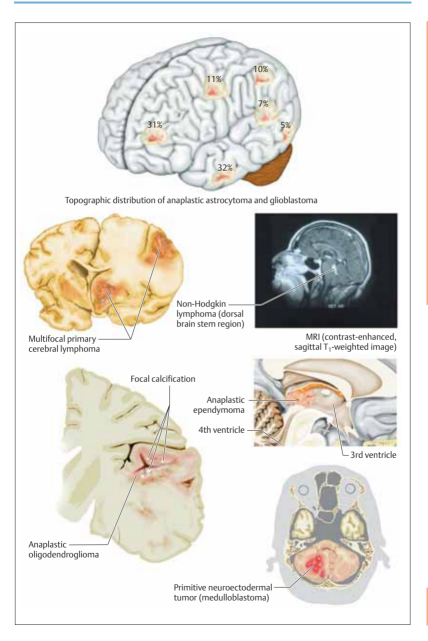
Primitive Neuroectodermal Tumor (PNET; WHO grade IV)

A PNET is a highly malignant embryonal tumor of the CNS that mainly arises in children. PNETs arising in the cerebellum, called *medulloblastomas*, are most commonly found in the vermis; they tend to metastasize to the leptomeninges and subarachnoid space (*drop metastasis*). The primary tumor and its metastases are best seen on MRI; they may appear in CT scans as areas of hyperdensity.

Primary Cerebral Sarcoma (WHO grade IV)

The very rare tumors in this group, including meningeal sarcoma, fibrosarcoma, chondrosarcoma, rhabdomyosarcoma, and malignant fibrous histiocytoma, all tend to recur locally and only rarely metastasize.

Brain Tumors



Metastatic Disease

Metastases spread to the nervous system through the bloodstream (cerebral, spinal, and leptomeningeal metastases), lymphatic vessels (metastases to the PNS), and cerebrospinal fluid (so-called drop metastases in the spinal subarachnoid space). Aside from direct metastatic involvement, the nervous system can also be affected by local tumor infiltration (e.g., of the brachial plexus by a Pancoast tumor), by external compression (e.g., of the spinal cord by a vertebral tumor, or of a peripheral nerve by a tumor-infiltrated lymph node), or by perineural infiltration (e.g. melanoma or salivary gland carcinoma). Only a small fraction of proliferating tumor cells are capable of metastasizing; thus, the biological behavior and drug response of metastasizing cells may differ from that of the primary tumor. Angiogenesis is essential for tumor growth and metastasis. Local invasion of surrounding tissue by the primary tumor makes it possible for tumor cells to break off and metastasize by way of the lymphatic vessels, veins, and arteries. Metastatic cells often settle in a vascular bed just downstream from the site of the primary tumor, thus (depending on its location) in the lungs, liver, or vertebral bodies. The nervous system may become involved thereafter in a second phase of metastasis (cascade hypothesis), or else directly, in which case the metastasizing cells must have passed through the intervening capillary bed without settling in it. Metastases may also bypass the lungs through a patent foramen ovale (paradoxical embolism).

Intracranial Metastases

Of all intracranial metastases, 85% are supratentorial, 15% infratentorial. The primary process in men is usually a tumor of the lung, gastrointestinal tract, or urogenital system, in women a tumor of the breast, lung, or gastrointestinal tract. Prostate, uterine, and gastrointestinal tumors metastasize preferentially to the cerebellum. The clinical manifestations of intracranial metastases are usually due to their local mass effect and surrounding cerebral edema. Brain metastases of melanoma, choriocarcinoma, and testicular cancer tend to produce hemorrhages. Metastases to the calvaria are usually asymptomatic. Skull base metastases cause pain and cranial nerve deficits. Dural-based metastases may compress or infiltrate the adjacent brain tissue, or exude fluid containing malignant cells into the subdural space. Pituitary metastases (mainly of breast cancer) cause endocrine dysfunction and cranial nerve deficits.

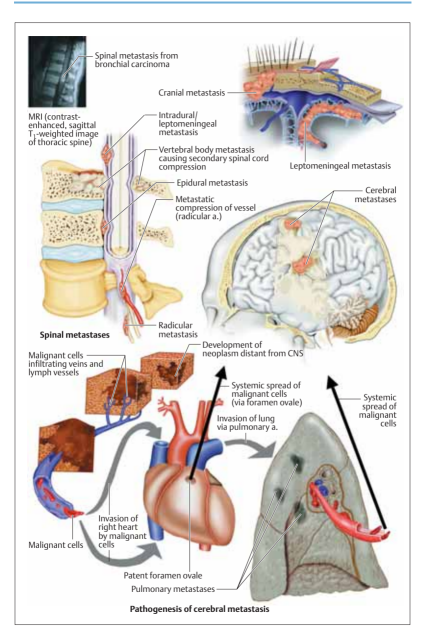
Spinal Metastases

The clinical manifestations of vertebral metastases, including vertebral or radicular pain, paraparesis/paraplegia, and gait ataxia, are mainly due to epidural mass effect. The bone marrow itself being insensitive to pain, pain arises only when the tumor compresses the periosteum, paravertebral soft tissue, nerve roots, or spinal cord. Spinal instability and pathological fractures cause additional pain. Pain in the spine may be the first sign of spinal metastasis. Subarachnoid and intramedullary metastases are rare (<5%).

Leptomeningeal Metastases (Neoplastic Meningeosis, "Carcinomatous Meningitis")

Seeding of the meninges may be diffuse or multifocal. Meningeal metastases may spread into the adjacent brain or spinal cord tissue, cranial nerves, or spinal nerves. *Cerebral* leptomeningeal involvement produces headache, gait ataxia, memory impairment, epileptic seizures, and cranial nerve deficits (e.g., facial nerve palsy, hearing loss, vertigo, diplopia, and loss of vision). *Spinal* involvement produces neck or back pain, radicular pain, paresthesia, paraparesis, and atony of the bowel and bladder.

Metastases



Classification and Treatment

As the treatment and prognosis of brain tumors depend on their histological type and degree of malignancy, the first step of management is tissue diagnosis (see Table 31, p. 377). The subsequent clinical course may differ from that predicted by the histological grade because of "sampling error" (i.e., biopsy of an unrepresentative portion of the tumor). Other factors influencing prognosis include age, the completeness of surgical resection, the preoperative and postoperative neurological findings, tumor progression, and the site of the tumor.

■ Incidence (adapted from Lantos et al., 1997)

The most common primary intracranial tumors in patients under 20 years of age are medulloblastoma, pilocytic astrocytoma, ependymoma, and astrocytoma (WHO grade II); from age 20 to age 45, astrocytoma (WHO grade II), oligodendroglioma, acoustic neuroma (schwannoma), and ependymoma; over age 45, glioblastoma, meningioma, acoustic neuroma, and oligodendroglioma. The overall incidence of pituitary tumors (including pituitary metastases), craniopharyngioma, and intracranial lymphoma and sarcoma is low.

■ Severity (Table 32, p. 378)

The Karnofsky scale (Karnofsky et al., 1951) is a commonly used measure of neurological disability, e. g., due to a brain tumor. Its use permits a standardized assessment of clinical course.

Treatment

The initial treatment is often neurosurgical, with the objective of removing the tumor as completely as possible without causing a severe or permanent neurological deficit. The resection can often be no more than subtotal because of the proximity of the tumor to eloquent brain areas or the lack of a distinct boundary between the tumor and the surrounding tissue. The overall treatment plan is usually a combination of different treatment modalities, chosen with consideration of the patient's general condition and the location, extent, and degree of malignity of the tumor.

Symptomatic treatment. *Edema*: The antiedematous action of glucocorticosteroids takes effect several hours after they are administered; thus, acute intracranial hypertension must be treated with an intravenously given osmotic agent (20% mannitol). Glycerol can be given orally to lower the corticosteroid dose in chronic therapy. Antiepileptic drugs (e.g., phenytoin or carbamazepine) are indicated if the patient has already had one or more seizures, or else prophylactically in patients with rapidly growing tumors and in the acute postoperative setting. Pain often requires treatment (headache, painful neoplastic meningeosis, painful local tumor invasion; cf. WHO staged treatment scheme for cancer-related pain). Restlessness: treatment of cerebral edema, psychotropic drugs (levomepromazine, melperone, chlorprothixene). Antithrombotic prophylaxis: Subcutaneous heparin.

Grade I tumors. Some benign tumors, such as those discovered incidentally, can simply be observed—for example, with MRI scans repeated every 6 months—but most should be surgically resected, as a total resection is usually curative. Residual tumor after surgery can often be treated radiosurgically (if indicated by the histological diagnosis). Pituitary tumors and craniopharyngiomas can cause endocrine disturbances. Meningiomas and craniopharyngiomas rarely recur after (total) resection.

Grade II tumors. Five-year survival rate is 50-80%. Complete surgical resection of grade II tumors can be curative. As these tumors grow slowly, they are often less aggressively resected than malignant tumors, so as not to produce a neurological deficit (partial resection, later resection of regrown tumor if necessary). Observation with serial MRI rather than surgical resection may be an appropriate option in some patients after the diagnosis has been established by stereotactic biopsy; surgery and/or radiotherapy will be needed later in case of clinical or radiological progression. Chemotherapy is indicated for unresectable (or no longer resectable) tumors, or after failure of radiotherapy

Grade III tumors. Patients with grade III tumors survive a median of 2 years from the time of diagnosis with the best current treatment involving multiple modalities (surgery, radiotherapy, chemotherapy). Many patients, however, live considerably longer. There are still inadequate data on the potential efficacy of chemotherapy against malignant forms of meningioma, plexus papilloma, pineocytoma, schwannoma, hemangiopericytoma, and pituitary adenoma.

Grade IV tumors. Patients with grade IV tumors survive a median of ca. 10 months from diagnosis even with the best current multimodality treatment (surgery, radiotherapy, chemotherapy). The 5-year survival rate of patients with glioblastoma is no more than 5%. PNET (including medulloblastoma) and primary cerebral lymphoma have median survival times of a few years.

Cerebral metastases: Solitary, surgically accessible metastases are resected as long as there is no acute progression of the underlying malignant disease, or for tissue diagnosis if the primary tumor is of unknown type. Solitary metastases of diameter less than 3 cm can also be treated with local radiotherapy, in one of two forms: interstitial radiotherapy with surgically implanted radioactive material (brachytherapy), or stereotactic radiosurgery. The latter is a closed technique, requiring no incision, employing multiple radioactive cobalt sources (as in the Gamma Knife and X-Knife) or a linear accelerator. Solitary or multiple brain metastases in the setting of progressive primary disease are generally treated with whole-brain irradiation. Chemotherapy is indicated for tumors of known responsiveness to chemotherapy in patients whose general condition is satisfactory. Metastatic small-cell lung cancer, primary CNS lymphoma, and germ cell tumors are treated with radiotherapy or chemotherapy rather than surgery.

Spinal metastases: Resection and radiotherapy for localized tumors; radiotherapy alone for diffuse metastatic disease.

Leptomeningeal metastases: Chemotherapy (systemic, intrathecal, or intraventricular); irradiation of neuraxis.

Aftercare

Follow-up examinations are scheduled at shorter or longer intervals depending on the degree of malignity of the neoplasm and on the outcome of initial management (usually involving some combination of surgery, radiotherapy, and chemotherapy), with adjustment for individual factors and for any complications that may be encountered in the further course of the disease. A single CT or MRI scan 3 months postoperatively may suffice for the patient with a completely resected, benign tumor, while patients with malignant tumors should be followed up by examination every 6 weeks and neuroimaging every 3 months, at least initially. Later visits can be less frequent if the tumor does not recur

Traumatic Brain Injury (TBI)

The outcome of traumatic brain injury depends on the type and extent of the acute (primary) injury and its secondary and late sequelae.

Direct/indirect history. A history of the precipitating event and of the patient's condition at the scene should be obtained from the patient (if possible), or from an eyewitness, or both. Vomiting or an epileptic seizure in the acute aftermath of the event should be noted. Also important are the past medical history, current medications (particularly anticoagulants), and any history of alcoholism or drug abuse.

Physical examination. *General:* Open wounds, fractures, bruises, bleeding or clear discharge from the nose or ear. *Neurological:* Respiration, circulation, pupils, motor function, other focal signs.

Diagnostic studies. *Laboratory:* Blood count, coagulation, electrolytes, blood glucose, urea, creatinine, serum osmolality, blood alcohol, drug levels in urine, pregnancy testing if indicated.

Essential radiological studies: Head CT with brain and bone windows is mandatory in all cases unless the neurological examination is completely normal. A cervical spine series from C1 to C7 is needed to rule out associated cervical injury. Plain films of the skull are generally unnecessary if CT is performed.

Additional studies, as indicated: Cranial or spinal MRI or MR angiography, EEG, Doppler ultrasonography, evoked potentials.

In multiorgan trauma: Blood should be typed and cross-matched and several units should be kept ready for transfusion as needed. Physical examination and ancillary studies for any fractures, abdominal bleeding, pulmonary injury.

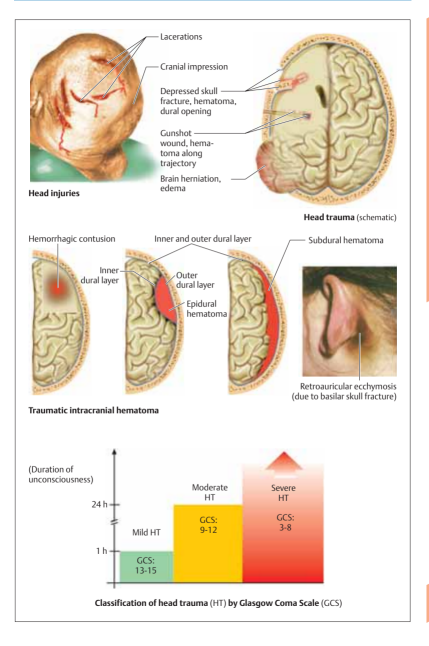
Primary Injury

The primary injury affects different parts of the skull and brain depending on the precipitating event. The traumatic lesion may be focal (hematoma, contusion, infarct, localized edema) or diffuse (hypoxic injury, subarachnoid hemorrhage, generalized edema). The worse the injury, the more severe the impairment of consciousness (pp. 116 ff). The clinical assessment of impairment of consciousness is described on pp. 378 f (Tables 33 and 34).

Region	Type of Injury	
Scalp	Cephalhematoma (neonates), laceration, scalping injury	
Skull	 Fracture mechanism: Bending fracture (caused by blows to the head, etc.), burst fracture (caused by broad skull compression) Fracture type: Linear fracture (fissure, fissured fracture, separation of cranial sutures), impression fracture, fracture with multiple fragments, puncture fracture, growing skull fracture (in children only) Fracture site: Convexity (calvaria), base of skull Basilar skull fracture: Frontobasal (bilateral periorbital hematoma ("raccoon sign"), bleeding from nose/mouth, CSF rhinorrhea, facial nerve palsy) Fracial skull fracture: LeFort I–III midface fracture; orbital base fracture 	
Dura mater	Open head trauma ¹ , CSF leak, pneumocephalus, pneumatocele	
Blood vessels	Acute epidural, subdural, subarachnoid or intraparenchymal hemorrhage; carotid- cavernous sinus fistula; arterial dissection	
Brain ²	 Contusion Diffuse axonal injury (clinical features: coma, autonomic dysfunction, decortication or decerebration, no focal lesion on CT or MRI) Penetrating (open or closed³) injury, or perforating (open) injury Brainstem injury 	

1 Wound with open dura and exposure of brain (definition). 2 Excluding cranial nerve lesions. 3 Without dural penetration (definition).

Trauma



Secondary Sequelae of TBI

Type of Sequela	Location/Syndrome	Special Features
Neurological		
Hematoma	⇔ Epidural	⇒ Lucid interval ¹ , immediate uncon- sciousness, or progressive deterioration of consciousness
	⇔ Subdural	May be asymptomatic at first, with progressive decline of consciousness
	⇔ Subarachnoid	⇔ Meningism
	\Rightarrow Intraparenchymal	⇒ Intracranial hypertension, ûfocal signs; often a severe injury
 Intracranial hypertension 	 Cerebral edema, hydrocephalus, massive hematoma 	• See p. 162. Risk of herniation
 Ischemia 	 Vasospasm, arterial dissection, fat embolism 	Acute focal signs
Epileptic seizure	Focal or generalized	Common in focal injury
Infection	CSF leak, open head injury	 Recurrent meningitis, encephalitis, empyema, abscess, ventriculitis
Amnesia	Anterograde/retrograde	• See p. 134
General		
 Hypotension, hypoxia, anemia 	Shock, respiratory failure	 Multiple trauma, pneumothorax or hemothorax, pericardial tamponade, blood loss, coagulopathy
• Fever, meningitis	Infection	• Pneumonia, sepsis, CSF leak
• Fluid imbalance	Hypothalamic lesion	• Diabetes insipidus ² , SIADH ³

1 Patient immediately loses consciousness \square awakens and appears normal for a few hours \square again loses consciousness. **2** Polyuria, polydipsia, nocturia, serum osmolality > 295 mOsm/kg, \square urine osmolality. **3** Syndrome of inappropriate secretion of ADH: euvolemia, serum osmolality < 275 mOsm/kg, excessively concentrated urine (urine osmolality > 100 mOsm/kg), \square urinary sodium despite normal salt/water intake; absence of adrenal, thyroid, pituitary and renal dysfunction.

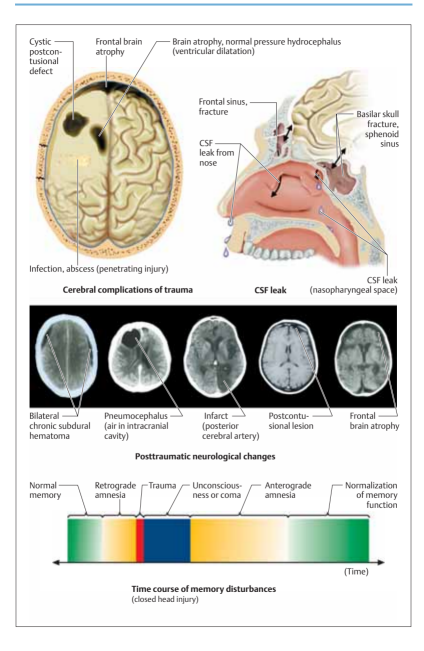
For overview of late complications of head trauma, see p. 379 (Table 35).

Prognosis

Head trauma causes physical impairment and behavioral abnormalities whose severity is correlated with that of the initial injury.

Severity ¹	Prognosis	
Mild	Posttraumatic syndrome resolves within 1 year in 85–90% of patients. The re- maining 10–15% develop a chronic posttraumatic syndrome	
Moderate	The symptoms and signs resolve more slowly and less completely than those of mild head injury. The prognosis appears to be worse for focal than for diffuse injuries. Reliable data on the long-term prognosis are not available	
Severe	Age-dependent mortality ranges from 30% to 80%. Younger patients have a bet- ter prognosis than older patients. Late behavioral changes (impairment of memory and concentration, abnormal affect, personality changes)	

1 For severity of head trauma, cf. pp. 378 f, Tables 33 and 34.



269

Posttraumatic Headache

Posttraumatic headache may be acute (<8 weeks after head trauma) or chronic (>8 weeks). The duration and intensity of the headache are not correlated with the severity of the precipitating head trauma. It can be focal or diffuse, continuous or episodic. It often worsens with physical exertion, mental stress, and tension and improves with rest and stress avoidance. Its type and extent are highly variable. If the headache gradually increases in severity, or if a new neurological deficit arises, further studies should be performed to exclude a late posttraumatic complication, such as chronic subdural hematoma (p. 379).

Pathogenesis of Traumatic Brain Injury

Direct blunt or penetrating injuries of the head and acceleration/deceleration injuries can damage the scalp, skull, meninges, cerebral vasculature, ventricular system, and brain parenchyma. The term primary injury refers to the initial mechanical damage to these tissues. Traumatized brain tissue is more sensitive to physiological changes than nontraumatized tissue. Secondary injury is caused by cellular dysfunction due to focal or global changes in cerebral blood flow and metabolism. Mechanisms involved in secondary injury include disruption of the blood-brain barrier, hypoxia, neurochemical changes (increased concentrations of acetylcholine, norepinephrine, dopamine, epinephrine, magnesium, calcium, and excitatory amino acids such as glutamate), cytotoxic processes (production of free radicals and of calcium-activated proteases and lipases), and inflammatory responses (edema, influx of leukocytes and macrophages, cytokine release). Epidural hematoma. Bleeding into the epidural space (pp. 6, 267) due to detachment of the outer dural sheath from the skull and rupture of a meningeal artery (usually the middle meningeal artery, torn by a linear fracture of the temporal bone). Epidural hematoma is less frequently of venous origin (usually due to tearing of a venous sinus by a skull fracture).

Subdural hematoma. Bleeding into the subdural space (pp. 6, 176, 267) because of disruption of larger bridging veins; often accompanied by focal contusion of the underlying brain. Frequently located in the temporal region.

Intracerebral hematoma. Bleeding into the tissue of the brain (intraparenchymal hematoma) under the site of impact, on the opposite side (*contre-coup*), or in the ventricular system (intraventricular hemorrhage) (pp. 176, 267).

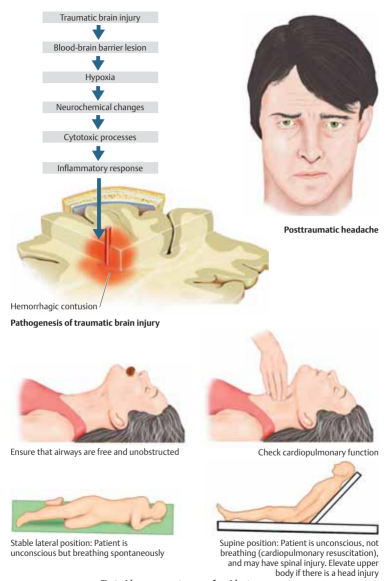
Subarachnoid hemorrhage. Rupture of pial vessels.

Treatment

At the scene of the accident. The scene should be secured to prevent further injury to the injured person, bystanders, or rescuers. First aid: Evaluation and clearing of the airway; cardiopulmonary resuscitation (CPR) if necessary. Immobilization of the cervical spine with a hard collar. Recognition and treatment of hemodynamic instability (keep systolic blood pressure above 120 mmHg), fluid administration as needed ("small volume resuscitation" with hyperoncotic-hypertonic solutions'). Dressing of wounds, sedation if necessary to reduce agitation, elevation of the upper body to 30°. Documentation: Time and nature of accident, general and neurological findings, drugs given. Transport: Cardiorespiratory monitoring.

In the hospital. Systematic assessment and treatment by organ system, with documentation of all measures taken. Cardiorespiratory monitoring: monitoring of blood gases and blood pressure (cerebral perfusion pressure >60-70 mmHg, p. 162). Respiratory system: Supplementary oxygen, intubation, and ventilation as needed. Cardiovascular system: Central venous access, administration of fluids and pressors as needed. Treatment of fever or hyperthermia. Administration of anticonvulsants as needed. Evaluation of tetanus vaccination status. Immediate neurosurgical consultation regarding the possible need for surgery. Treatment of intracranial hypertension: Sedatives, analgesics; if ICP (p. 162) is above 20-25 mmHg, osmotherapy with 20% mannitol, bolus of 0.35 mg/kg over 10-15 minutes, repeated every 4-8 hours as needed: barbiturate coma (thiopental); decompressive bifrontal craniectomy may be indicated in refractory cerebral edema.

Trauma



271

First-aid measures at scene of accident

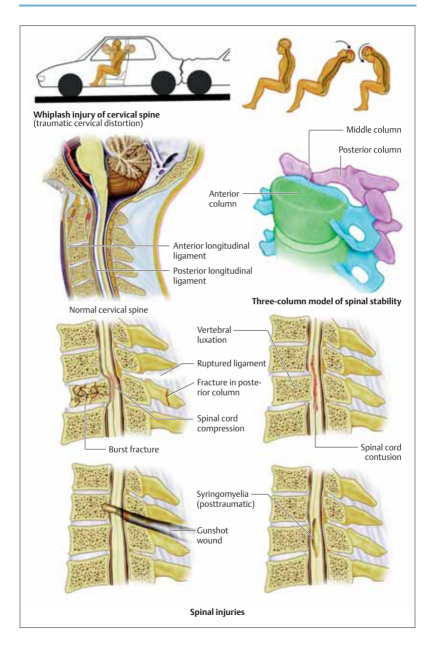
Spinal Trauma

Spinal injury can involve the vertebrae, ligaments, intervertebral disks, blood vessels, muscles, nerve roots, and spinal cord. The spinal cord and spinal nerve roots may be directly injured (e. g. by gunshot or stab wounds) or secondarily affected by compression (bone fragments), hyperextension (spinal instability), and vascular lesions (ischemia, hemorrhage). *Diagnosis*: Bone injuries can be identified by radiography and/or CT; spinal cord lesions (hemorrhage, contusion, edema, transection) and softtissue lesions (hematoma, edema, arterial dissection) are best seen on MRI.

Cervical spine distortion (whiplash injury). Indirect spinal trauma (head-on or rear-end collision) leads to sudden passive retroflexion and subsequent anteflexion of the neck. The forces acting on the spine (acceleration, deceleration, rotation, traction) can produce both cervical spine injuries (spinal cord, nerve roots, retropharyngeal space, bones, ligaments, joints, intervertebral disks, blood vessels) and cranial injuries (brain, eyes, temporomandibular joint). There may be an interval of 4-48 hours until symptoms develop, rarely longer (asymptomatic period). Symptoms and signs: Pain in the head, neck, and shoulders, neck stiffness, and vertigo may be accompanied by forgetfulness, poor concentration, insomnia, and lethargy. The symptoms usually resolve within 3-12 months but persist for longer periods in 15-20% of patients, for unknown reasons. Severity classification: Grade I = no neurological deficit or radiological abnormality, grade II = neurological deficit without radiological abnormality, grade III = neurological deficit and radiological abnormality.

Vertebral fracture. It must be determined whether the fracture is stable or unstable; if it is unstable, any movement can cause (further) damage to the spinal cord and nerve roots. Thus, all patients who may have vertebral fractures must be transported in a stabilized supine position, with the head in a neutral position (e.g., on a vacuum mattress). Repositioning the patient manually with the "collar splint grip," "paddle grip," or "bridge grip" should be avoided if possible. In the assessment of stability, it is useful to consider the spinal column and intervertebral disks as composed of three columns. Involvement of only one column = stable injury; two columns = potentially unstable; three columns = unstable. For details, see p. 380 (Table 36).

Trauma to nerve roots and brachial plexus. Nerve root lesions usually involve the ventral roots, and thus usually produce a motor rather than sensory deficit. Nerve root avulsion may be suspected on the basis of (multi)radicular findings and/or Horner syndrome and can be confirmed by myelography (empty root sleeves, bulging of the subarachnoid space) or MRI. Downward or backward traction on the shoulder and arm (as in a motorcycle accident) can produce severe brachial plexus injuries accompanied by nerve root avulsion. Brachial plexus lesions can also be caused by improper patient positioning during general anesthesia, intense supraclavicular pressure (backpack paralysis), or local trauma (stab or gunshot wound, bone fragments, contusion, avulsion). These injuries more commonly affect the upper portion of the brachial plexus (pp. 34, 321).



Spinal Cord Trauma

Open spinal cord trauma, by definition, involves penetration of the dura mater by a stab wound, gunshot wound, bone fragment, or severely dislocated vertebra. *Closed* spinal cord trauma (with dura intact) is the indirect effect of a non-penetrating injury. The result may be a complete or incomplete spinal cord transection syndrome (p. 48; Table 37, p. 380).

Acute stage (spinal shock). The acute manifestations of spinal cord transection syndrome are seen below the level of the injury and include the total loss of voluntary and reflex *motor function* (flaccid paraplegia or quadriplegia, areflexia) and *sensation*, and *autonomic dysfunction* (urinary retention \Rightarrow overflow incontinence, intestinal atony \Rightarrow paralytic bowel obstruction, anhidrosis \Rightarrow hyperthermia, cardiovascular dysfunction \Rightarrow orthostatic hypotension, cardiac arrhythmia, paroxysmal hypertension). Patients are usually stable enough to begin rehabilitation in 3–6 weeks (rehabilitation stage, see below). For acute treatment, see p. 380 (Table 38).

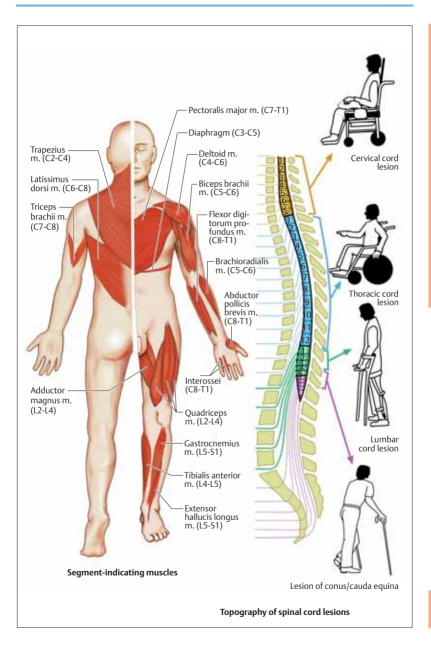
Level ¹	Motor Deficit	Sensory Deficit ²	Autonomic Deficit ³
C1-C3 ⁴	Quadriplegia, neck muscle paresis, spasticity, respira- tory paralysis	Sensory level at back of head/edge of lower jaw; pain in back of head, neck, and shoulders	Voluntary control of bladder, bowel, and sexual function replaced by reflex control; Horner syndrome
C4-C5	Quadriplegia, diaphragmatic breathing	Sensory level at clavicle/ shoulder	Same as above
C6–C8 ⁵	Quadriplegia, spasticity, flac- cid arm paresis, diaphrag- matic breathing	Sensory level at upper chest wall/back; arms involved, shoulders spared	Same as above
T1-T5	Paraplegia, diminished respi- ratory volume	Sensory loss from inner sur- face of lower arm, upper chest wall, back region downward	Voluntary control of bladder, bowel, and sexual function replaced by reflex control
T5-T10	Paraplegia, spasticity	Sensory level on chest wall and back corresponding to level of spinal cord injury	Same as above
T11-L3	Flaccid paraplegia	Sensory loss from groin/ven- tral thigh downward, de- pending on level of injury	Same as above
L4-S2 ⁶	Distal flaccid paraplegia	Sensory loss at shin/dorsum of foot/posterior thigh downward, depending on level of injury	Flaccid paralysis of bladder and bowel, loss of erectile function
S3-S57	No motor deficit	Sensory loss in perianal re- gion and inner thigh	Flaccid paralysis of bladder and bowel, loss of erectile function

Rehabilitation stage. The neurological deficits depend on the level of the lesion.

1 Spinal cord level (not the same as vertebral level). 2 See p. 32 ff. 3 Disturbance of bladder, bowel, rectal, and erectile function, sweating, and blood pressure regulation; p. 140 ff. 4 High cervical cord lesion. 5 Low cervical cord lesion. 6 Epiconus. 7 Conus medullaris.

Chronic stage—late sequelae. Persistence of neurological deficits; assorted complications including venous thrombosis, pulmonary embolism, respiratory insufficiency, bowel obstruction, urinary tract infections, sexual dysfunction, cardiovascular disturbances, spasticity, chronic pain, bed sores, heterotopic ossification, and syringomyelia.

Trauma



Signs of Cerebellar Dysfunction

Loss of coordination and balance. Ataxia is uncoordinated, irregular, and poorly articulated movement (dyssynergy). The typical patient sways while sitting (truncal ataxia) or standing (postural ataxia), undershoots or overshoots an intended target of movement (dysmetria = hypometria or hypermetria), and walks with quick, irregular steps in an unsteady, swaying, broadbased gait reminiscent of alcohol intoxication (gait ataxia, p. 54). Pointing tests are used to detect dysmetria, incoordination, and tremor that is worst as a movement approaches its target (intention tremor); the finger-nose, finger-finger, and heel-knee-shin tests should be carried out with the eyes open and closed. Bárány's pointing test: The patient is asked to close his or her eyes, touch the doctor's finger with his or her own index finger, then lower and raise the still outstretched arm and touch the doctor's finger again; the patient's finger deviates laterally from the target, and the direction of deviation is toward the side of the lesion. Unsteadiness of stance of cerebellar origin, which may be so severe as to make standing impossible (astasia), is not influenced by opening or closing the eyes (Romberg sign) and differs in this respect from spinal (sensory) ataxia. Stepping in place for 30-60 seconds with the eyes closed causes the body to turn to the side of the lesion. Patients with mild ataxia find it difficult or impossible to walk a straight line (abasia; detected by heel-totoe walking, tandem gait). The patient may be unable to perform rapid alternating movements (dysdiadochokinesia). The handwriting is enlarged (macrographia), coarse, and shaky, and the patient's drawing of parallel lines or a spiral is unsatisfactory.

Dysarthria. The patient's speech (p. 130) is slow, unclear (babbling, slurred), and monotonous (*dysarthrophonia*), and possibly also discontinuous (choppy, faltering, or scanning speech). There is poor coordination of breathing with the flow of speech, resulting in a sudden transition from soft to loud speech (explosive speech).

Oculomotor disturbances. Gaze-evoked nystagmus is a frequent finding in cerebellar disease. Voluntary saccades are too short or too long (*ocular dysmetria*) and are therefore followed by afterbeats. Slow pursuit movements are jerky (saccadic). Patients are frequently unable to suppress the vestibulo-ocular reflex (p. 26), i.e., the normal visual suppression of nystagmus is impaired. The result is impaired visual fixation on turning of the head.

Muscle tone. Decreased muscle tone is mainly found in patients with acute unilateral lesions of the cerebellum. The examiner can detect it by passively swinging or shaking the patient's limbs, or by testing for the *rebound phenome-non*. The patient is asked to extend the arms with the eyes closed (*posture test*) and the examiner lightly taps on one wrist, causing deflection of the arm. The rebound movement undershoots or overshoots the original arm position. Alternatively, the patient can be asked to flex the elbow against resistance. When the examiner suddenly releases the resistance, the affected arm rebounds unchecked.

Topography of Cerebellar Lesions

Lesions of the cerebellum and its afferent and efferent connections (p. 54) produce characteristic signs of cerebellar disease. Expanding lesions may go on to produce further, extracerebellar deficits (e.g., cranial nerve palsies, hemiparesis, sensory loss).

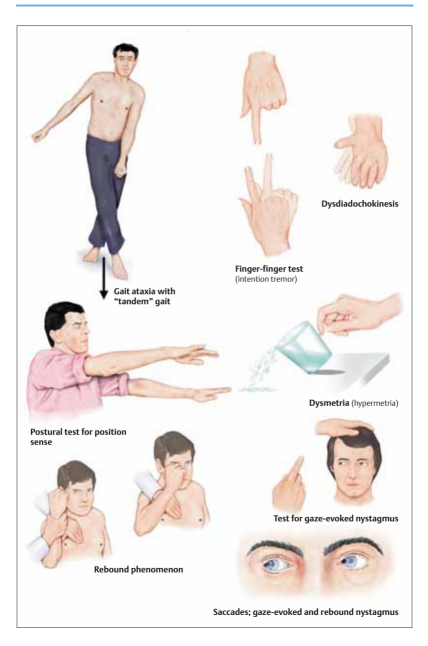
Special Diagnostic Studies

The diagnostic studies to be obtained depend on the clinical findings (to be described below) and may include imaging studies (MRI, CT), neurophysiological studies (nerve conduction studies, electromyography), ECG, pathological studies (of tissue, blood, CSF, bone marrow, muscle, or nerve biopsy specimens), and/or ophthalmological consultation (optic nerve atrophy, Kayser–Fleischer ring, tapetoretinal degeneration).

Idiopathic Cerebellar Ataxia (IDCA)

This group of disorders includes various forms of nonfamilial cerebellar ataxia of unknown cause with onset in adulthood (generally age 25 years or older). IDCA occurs as an isolated disturbance or as a component of multiple system atrophy (MSA; p. 302).

Cerebellar Diseases



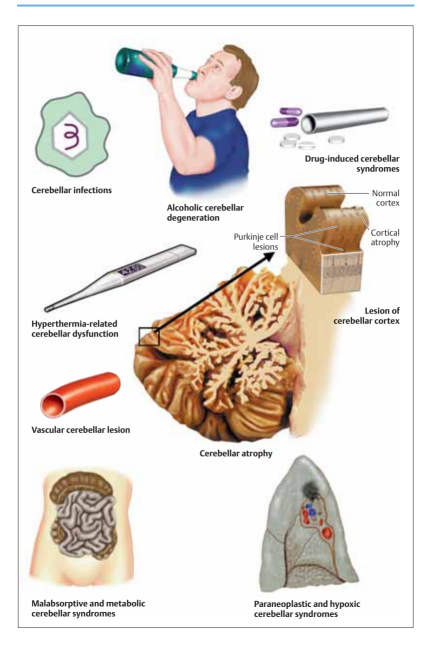
277

Acquired Cerebellar Syndromes

Onset	Etiology	Symptoms and Signs
Acute (minutes to hours)	• Infection ¹	 Viral infection: varicella-zoster virus, Epstein–Barr virus, rubella, mumps, influenza, parainfluenza, echovirus, coxsackievirus, cytomegalovirus, FSME, herpes simplex virus. Children are more commonly affected than adults. Special type: opsoclonus-ataxia syndrome². Abscess Miller Fisher syndrome (ataxia, ophthalmoplegia, areflexia; p. 395)
	• Vascular	 Brainstem signs (pp. 70 ff., 170) predominate Infarcts can be differentiated from hemorrhages by imaging studies Early treatment, often neurosurgical, may be needed to prevent rapid development of life-threatening complications (p. 174 f)
	Toxic	Alcohol, barbiturates, phenytoin, lithium
Subacute (days to weeks)	 Tumor³ Paraneoplastic⁴ 	 Occipital pain (radiating to forehead, nuchal region, and shoulders), recurrent vomiting, stiff neck, vertigo, truncal ataxia; obstructive hydrocephalus Cerebellar dysfunction may appear months or years before the tumor is discovered. Anti-Purkinje-cell antibodies are pre- sent in the serum and CSF of patients with neuron loss
	ToxicOther	 Alcohol Medications (anticonvulsants, e. g., phenytoin; lithium, 5-fluorouracil, cytosine arabinoside) Heavy metals (mercury, thallium, lead) Solvents (toluene, carbon tetrachloride) Hypoxia, heat stroke, hyperthermia
Chronic (months to years)	• Infection	 Progressive rubella panencephalitis (very rare complication of congenital rubella infection in boys; onset at age 8 to 19 years; characterized by ataxia, dementia, spasticity, and dysarthria) Creutzfeldt–Jakob disease (p. 252)
	VascularMetabolic	 Meningeal siderosis causes ataxia and partial or complete hearing loss (leptomeningeal deposition of hemosiderin in chronic subarachnoid hemorrhage ⇒ vascular malformations, oligodendroglioma, ependymoma of the cauda equina, postoperative occurrence) Hypothyroidism, malabsorption syndrome (vitamin E deficiency), thiamin deficiency (acute ⇒ Wernicke encephalopathy) Refsum disease⁵ (i)serum phytanic acid level, p. 332) Wilson disease⁵ (ataxia, tremor, dysarthrophonia, dysphagia,
Intermittent	• Metabolic ^s	 dystonia, behavioral disturbances, p. 307) Hereditary metabolic disorders in neonates, children, and juveniles (see also pp. 306 f, 386 f) Disorders of amino acid metabolism (hyperammonemia, Hartnup syndrome, maple syrup urine disease) Storage diseases (metachromatic leukodystrophy, neuronal ceroid lipofuscinosis, sialidosis, GM₂ gangliosidosis)

1 Partial listing; numerous infections can cause ataxia as part of the syndrome of encephalomyelitis. 2 Highfrequency bursts of saccades in all directions of gaze without an intersaccadic interval. 3 See p. 254 ff; cerebellar astrocytoma, medulloblastoma, ependymoma, hemangioblastoma (von Hippel-Lindau disease), meningioma of the cerebellopontine angle, metastases (lung cancer, breast cancer, melanoma). 4 Antibodies (p. 388) against Hu, Yo, TR, CV2, Ma1, CRD1, CRD2, Ma2, and mGluR1. 5 Genetic; listed here for differential diagnostic purposes.

Cerebellar Diseases



279

Hereditary Cerebellar Syndromes

Autosomal Recessive Cerebellar Syndromes (partial listing)

Syndrome	Symptoms and Signs	CL ¹ /Gene Product
Friedreich ataxia ^{2,7}	Usual manifestations: ● Progressive limb/gait ataxia A dge of onset < 30 years A reflexia in legs ● Neurophysiological evidence of sensory neuropathy Variable manifestations: Dysarthria, distal muscular atrophy/paresis (ca. 50%), pes cavus (ca. 50%), scoliosis, optic nerve atrophy (ca. 25%), nystagmus (ca. 20%), oculomotor disturbances (p. 276), hearing loss (ca. 10%), cardiomy- opathy (ca. 65%), diabetes mellitus (ca. 10%)	9q13, 9p23-p11/ frataxin <i>Mutation:</i> Extended GAA-trinucleotide repeat
Ataxia with vitamin E deficiency ⁷ (serum: the vitamin E, the cholesterol/ triglycerides)	 Onset in childhood or adulthood Gait ataxia Dysarthria Other symptoms similar to those of Friedreich ataxia 	8q13.1-q13.3/α-to- copherol transfer protein
Abetalipoprotein- emia ^{3,7} (p. 300)	Steatorrhea, other symptoms similar to those of Friedreich ataxia	4q24/triglyceride transfer protein
Ataxia-telangiec- tasia ^{4,7}	 Ataxia first seen when child learns to walk Choreoathetosis Oculomotor disturbances⁵ Oculocutaneous telangiectases Immunodeficiency (frequent infections) Increased risk of malignant tumors Elevated serum α-fetoprotein 	11q22.3/phosphatidyl- inositol-3'- kinase and rad3 ⁶

 Chromosome location (CL). 2 Classic form. 3 Bassen-Kornzweig syndrome; vitamin A and E deficiency, low cholesterol/ triglyceride levels, acanthocytosis. 4 Louis-Bar syndrome. 5 Oculomotor apraxia. 6 DNA repair kinase/cell cycle control; ataxia-telangiectasia-mutated (ATM) gene. 7 A direct gene test is available.

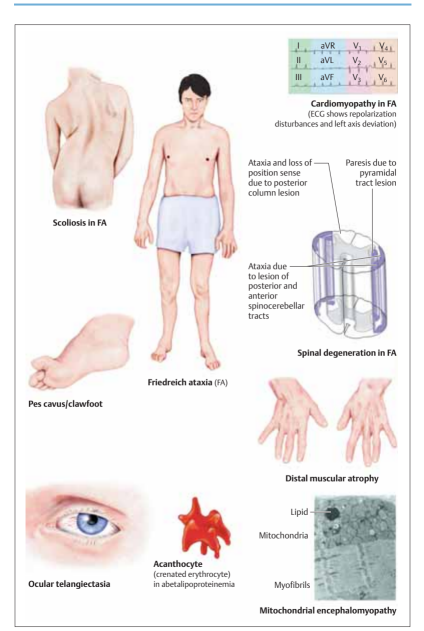
For mitochondrial syndromes with ataxia, see p. 403.

Autosomal Dominant Cerebellar Syndromes (partial listing)

Syndrome	Symptoms and Signs	CL/Gene Product
Autosomal domi- nant cerebellar ataxia (ADCA); spinocerebellar ataxia (SCA) ¹	 ADCA1: Ataxia, ophthalmoplegia, pyramidal/extrapyramidal disturbances (p. 44); SCA1⁵, SCA3^{2,5}, SCA3^{2,5}, SCA4, SCA8⁵, SCA12, SCA13, SCA17 ADCA2: Ataxia, retinopathy, SCA7⁵ ADCA3: Predominant cerebellar ataxia; SCA5, SCA6⁵, SCA10, SCA11, SCA12⁵, SCA14, SCA15, SCA16 	SCA1: 6p23/ataxin1 SCA2: 12q24/ataxin2 SCA3: 14q24.3-q31/ MJD1 protein SCA4: 16q22.1 SCA5: 11p11-q11 SCA6: 19p13/ac-1A calcium channel SCA7: 3p21.1-p12
Episodic ataxia (EA) ³	 EA1: Episodes of ataxia lasting seconds to minutes, 1 to 10 times daily; provoked by abrupt changes of position, emotional or physical stress, and caloric vestibular stimulation; myokymia in face and hands be- tween attacks; continuous spontaneous activity in resting EMG EA2: Episodes of ataxia lasting minutes to hours (rarely days) of varia- ble frequency (daily to yearly); headache, tinnitus, vertigo, ataxia, nausea, vomiting, nystagmus; induced by same stimuli as EA1; ataxia, nystagmus, and head tremor between attacks 	 12p13⁵/potassium channel (point mutation) 19p13⁵/voltage- gated calcium channel⁴ (point mutation)
Gerstmann– Sträussler– Scheinker syn- drome (p. 252)	Onset between the ages 40 and 50 years; presents with cerebellar ataxia; dysarthrophonia, dementia, nystagmus, rigor, visual disturbances, and hearing loss develop in the course of the disease	20pter-p12/P102L
Fatal familial in- somnia (p. 252)	Progressive insomnia, autonomic dysfunction (arterial hypertension, tachycardia, hyperthermia, hyperhidrosis), myoclonus, tremor, ataxia	20pter-p12/D178N

280

1 Definitive identification of the SCA types listed is possible only with molecular genetics tests (examples in right column, see OMIM for details). 2 Machado–Joseph disease (MJD). 3 Other forms: EA3 and EA4. 4 Other mutations of this gene are associated with SCA6 and familial hemiplegic migraine. 5 A direct genetic test is available.



Myelopathies

The clinical differential diagnosis of myelopathies is based on the level of the spinal cord lesion, the particular structures affected, and the temporal course of the disorder (p. 48, Table 39, p. 381).

Acute Myelopathies

Symptoms and signs develop within minutes, hours, or days.

Spinal Cord Trauma

(See p. 274)

Myelitis

Viral myelitis (p. 234 ff). Enteroviruses (poliovirus, coxsackievirus, echovirus), herpes zoster virus, varicella zoster virus, FSME, rabies, HTLV-1, HIV, Epstein–Barr virus, cytomegalovirus, herpes simplex virus, postvaccinial myelitis.

Nonviral myelitis (p. 222 ff). Mycoplasma, neuroborreliosis, abscess (epidural, intramedullary), tuberculosis, parasites (echinococcosis, cysticercosis, schistosomiasis), fungi, neurosyphilis, sarcoidosis, postinfectious myelitis, multiple sclerosis/neuromyelitis optica (Devic syndrome), acute necrotizing myelitis, connective tissue disease (vasculitis), paraneoplastic myelitis, subacute myelo-optic neuropathy (SMON), arachnoiditis (after surgical procedures, myelography, or intrathecal drug administration).

Vascular Syndromes (p. 22)

Anterior spinal artery syndrome. Segmental paresthesia and pain radiating in a bandlike distribution may precede the development of motor signs by minutes to hours. A flaccid paraparesis or quadriparesis (corticospinal tract, anterior horn) then ensues, along with a dissociated sensory loss from the level of the lesion downward (spinothalamic tract \Rightarrow impaired pain and temperature sensation, with intact perception of vibration and position) and urinary and fecal incontinence. Often only some of these signs are present.

Posterior spinal artery syndrome is rare and difficult to diagnose. It is characterized by pain in the spine, paresthesiae in the legs, a loss of position and vibration sense below the level of the lesion, and global anesthesia with segmental loss of deep tendon reflexes at the level of the lesion. Larger lesions cause paresis and sphincter dysfunction.

Sulcocommissural artery syndrome. Segmental pain at the level of the lesion, followed by flaccid paresis of ipsilateral arm/leg; loss of proprioception, position sense, and touch perception with contralateral dissociated sensory loss (Brown– Séquard syndrome). Sphincter dysfunction is rare.

Complete spinal infarction. Acute spinal cord transection syndrome with flaccid paraplegia or quadriplegia, sphincter dysfunction, and total sensory loss below the level of the lesion. Autonomic dysfunction may also occur (e.g., vasodilatation, pulmonary edema, intestinal atony, disordered thermoregulation). The cause is often an acute occlusion of the great radicular artery (of Adamkiewicz).

Central spinal infarction. Acute paraplegia, sensory loss, and sphincter paralysis.

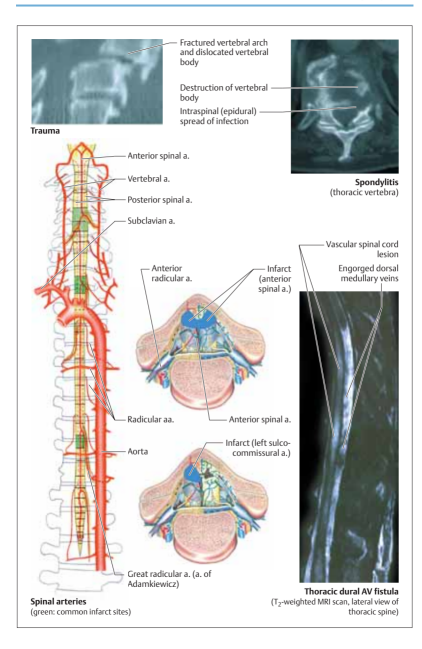
Claudication of spinal cord. Physical exercise (running, long walks) induces paresthesiae or paraparesis that resolves with rest and does not occur when the patient is lying down.

Cause: Exercise-related ischemia of the spinal cord due to a dural arteriovenous fistula or high-grade aortic stenosis (see also p. 284).

Dural/perimedullary arteriovenous (AV) fistula is an abnormal communication (shunt) between an artery and vein between the two layers of the dural mater. An arterial branch of a spinal artery feeds directly into a superficial spinal vein, which therefore contains arterial rather than venous blood, flowing in the opposite direction to normal. Paroxysmal stabbing pain and/or episodes of slowly progressing paraparesis and sensory loss separated by periods of remission occur in the early stage of the disorder, which usually affects men between the ages of 40 and 60. If the suspected diagnosis cannot be confirmed by MRI scans (because of low shunt volume), myelography may be helpful (⇒ dilated veins in the subarachnoid space).

Spinal hemorrhage can occur in epidural, subdural, subarachnoid, and intramedullary locations (intramedullary hemorrhage = hematomyelia). *Possible causes:* intradural/intramedullary AV malformation, cavernoma, tumor, aneurysm, trauma, lumbar puncture, and coagulopathy.

Myelopathies



Subacute and Chronic Myelopathies

Spinal cord syndromes (p. 282) may be subacute or chronic depending on their cause. The

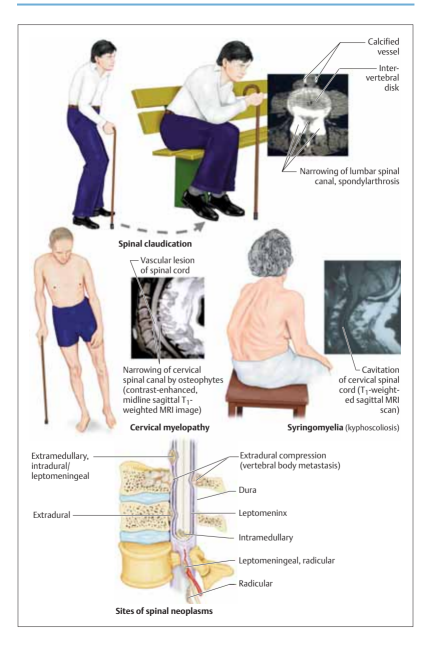
Mass Lesions

complete clinical picture may develop over days to weeks (subacute) or months to years (chronic). For myelopathies due to developmental disorders, see p. 288 ff.

Syndrome	Symptoms and Signs	Causes	Diagnosis/Treatment ¹
Cervical myelopathy	Progressive paraparesis or quadriparesis, spasticity, Iher- mitte's sign, reduced mobility of cervical spine; cervical radiculopathy may also occur	Spinal cord compres- sion ² by cervical spine lesions ³	MRI, CT, myelography; evoked potentials, EMG for radicular le- sions, plain radiograph of cervi- cal spine. <i>Treatment</i> : Surgery for progres- sive impairment or severe ste- nosis; otherwise, symptomatic treatment
Lumbar spinal stenosis ⁴ (intermittent claudication)	Early: Paresthesiae (sensation of heaviness) occur upon standing or walking (especially down stairs) and disappear with rest. <i>Late:</i> Only partial improvement of paresthesiae with rest; re- duced walking range	Compression of cauda equina by lumbar spine lesions ⁵	Diagnostic testing as above. Treatment: Surgery for severely decreased walking range or persistent symptoms; other- wise, analgesics and physi- otherapy (to strengthen trunk muscles)
Syringo- myelia ⁶	Pain, central cord deficits, kyphoscoliosis	Anomalous develop- ment of neural groove, obstruction of CSF flow, trauma, tumor	Diagnostic testing as above. <i>Treatment:</i> Surgery for progres- sive symptoms, especially pain ⁷
Neoplasm	Pain, sensory loss, segmental/ radicular paresis, Lhermitte's sign (cervical), incomplete or complete spinal cord transec- tion syndrome	Intramedullary: Ependymoma, glioma Extramedullary: Meningioma, neurofi- broma, vascular mal- formation Extradural: Metastasis, sarcoma	Diagnostic testing as above. <i>Treatment</i> : Surgery: radiother- apy if indicated; symptom con- trol with corticosteroids and analgesics

1 Principles of diagnosis and treatment. 2 Symptoms arise when sagittal diameter of spinal canal is in the range of 7–12 mm (normal 17–18 mm). 3 Primary spinal canal stenosis, disk protrusion/herniation, spinal degenerative disease (spondylosis, osteochondrosis), hyperextension of cervical spine (trauma, chiropractic maneuvers, dental procedures) in patient with cervical stenosis, Paget disease, or ossification of the posterior longitudinal ligament. 4 Not a myelopathy (the cauda equina is affected); mentioned here for differential diagnostic reasons. 5 Primary spinal canal stenosis, intervertebral disk protrusion/herniation, degenerative changes in spinal column. 6 Syringobulbia ⇒ pain (V), caudal cranial nerve lesions (VIII–XII), nystagmus. 7 Suboccipital decompression in Chiari malformation (p. 292), shunt ⇒ syringotomy.

Myelopathies



Non-Mass Lesions

Myelitis. See p. 282 for a listing of various infectious myelitides.

Subacute combined degeneration (SCD) appears in middle to old age, causing tingling and burning dysesthesiae in the limbs, gait unsteadiness, and abnormal fatigability. There may also be visual disturbances and depressive or psychotic symptoms accompanied by weight loss, glossopyrosis, and abdominal complaints. The neurological examination reveals a loss of position sensation (\Rightarrow spinal ataxia), spastic paraparesis, variable abnormalities of the deep tendon reflexes, and autonomic dysfunction (bladder, bowel, sexual dysfunction). Megalocytic anemia is usually present. The cause is vitamin B₁₂ deficiency, which may, in turn, be due to malabsorption, cachexia, or various medications. Folic acid deficiency produces a similar syndrome. The patient should be treated with parenteral cyanocobalamin or hydroxocobalamin as soon as possible. Neurological deficits can arise even if

the hematocrit and red blood cell count are normal.

Toxic myelopathy. Most patients initially present with polyneuropathy, developing clinically apparent myelopathy only in the later stages of disease. Common causes include solvent abuse ("glue sniffing"), a high dietary intake of gross peas (lathyrism; p. 304), and consumption of cooking oil adulterated with lubricant oil (triorthocresyl phosphate poisoning).

Hereditary. The clinically and genetically heterogeneous forms of *familial spastic paraplegia* (FSP; spinal paralysis = SPG, p. 384) become symptomatic either in the first decade of life or between the ages of 10 and 40. Progressive central paraparesis with spasticity arises either in isolation (uncomplicated SPG) or accompanied by variable neurological deficits (complicated SPG). Both types can be transmitted in an autosomal dominant, autosomal recessive, or X-linked inheritance pattern. *Spinal muscular atrophy*, see p. 304. *Adrenomyeloneuropathy*, see p. 384.

Method	Information Provided
Evoked potentials	SEP ² : conduction delay. MEP ³ : prolongation of CMT ⁴
Plain radiograph	Anomalies of spinal column or craniocervical junction, degenerative changes, fractures, lytic lesions, spondylolisthesis
СТ	Same as above, tumor, 3-D reconstruction
MRI	Tumor, myelitis, vascular myelopathy, (MR) myelography
Bone scan	Vertebral body lesions (trauma, neoplasm, inflammation, degeneration)
CSF analysis	Inflammatory, hemorrhagic (vascular), or neoplastic changes
Myelography ⁵	Position-dependent changes (dynamic spondylolisthesis), spinal stenosis, arachnoiditis, nerve root avulsion
Spinal angiography	Arteriovenous fistula/malformation, location of source of hemorrhage

Diagnostic Studies in Myelopathy¹

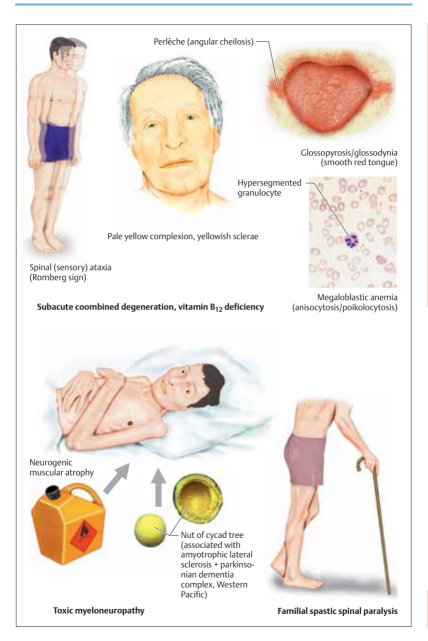
1 Urodynamic tests are used to evaluate bladder dysfunction (p. 156). 2 Somatosensory EP. 3 Motor EP. 4 Central motor conduction time (CMT). 5 Used when CT/MRI findings are ambiguous, in an emergency if CT and MRI are not available, or if position-dependent changes must be evaluated.

Treatment of Myelopathies (partial listing)

Cause	Treatment Measures
Myelitis	HSV/VZV1: acyclovir. Bacterial infection: antibiotics. Unknown pathogen: corticosteroids
Neoplasm ²	Surgical resection of tumor and stabilization of spinal column; radiotherapy; corti- costeroids; chemotherapy; hormonal therapy
Vascular lesion	AV malformation: Embolization, surgery. Ischemia/hematomyelia: symptomatic treat- ment, physiotherapy

1 HSV/VZV: herpes simplex virus/varicella-zoster virus. 2 Treatment depends on type and extent of neoplasm.

Myelopathies



Hereditary Diseases

Phenotype. The manner in which a hereditary disease expresses itself at a given moment in development (phenotype) is the product of both the individual's genetic makeup (genotype) and the environment in which development has taken place.

Inheritance. The human genome consists of 22 pairs of chromosomes (autosomes) and 2 sex chromosomes (either XX or XY). Individuals inherit half of their chromosomes from each parent. The chromosomes are made of DNA and bear the genes, sequences of nucleotide base pairs that encode the proteins of the body. Stretches of DNA that encode proteins are called exons; there are also intervening noncoding sequences, called introns. The inheritance pattern of hereditary diseases can be monogenic-the disease is due to a defect in a single (autosomal or X-chromosomal) gene, and is transmitted in a recessive or dominant manner in accordance with Mendel's laws: polygenic-the disease is due to defects in multiple genes; or multifactorial-the cause of disease is not exclusively genetic, and exogenous factors along with genetic factors determine its phenotype. Mitochondrial disorders are transmitted exclusively by maternal inheritance, as mitochondrial DNA is nonchromosomal and is inherited exclusively from the mother.

Mutation. Alleles are different forms of a gene. A gene mutation is a change in the DNA sequence of a gene and may involve a change in a single base pair (point mutation), the loss of one or more base pairs (deletion), the insertion of one or more base pairs, or unstable trinucleotide repeats. There are also genome mutations, which involve a change in the number of chromosomes, such as trisomy 21 (the cause of Down syndrome), as well as chromosome mutations, in which the chromosomal structure is altered. Mutations can occur either in the germ cells (germ-line mutation) or in the differentiated cells of the body (somatic mutation). Somatic mutations cause cancer, autoimmune diseases, and congenital anomalies.

Diagnosis. The diagnosis of hereditary diseases is by family history. Many monogenic diseases can be diagnosed by *direct* genotypic analysis (DNA sequencing). *Indirect* genotypic analysis, with investigation of the affected and nonaffected members of a single pedigree, is used in the diagnosis of disorders for which a gene locus is known but the responsible mutation(s) has not yet been determined.

Malformations and Developmental Anomalies (Table 40, p. 381)

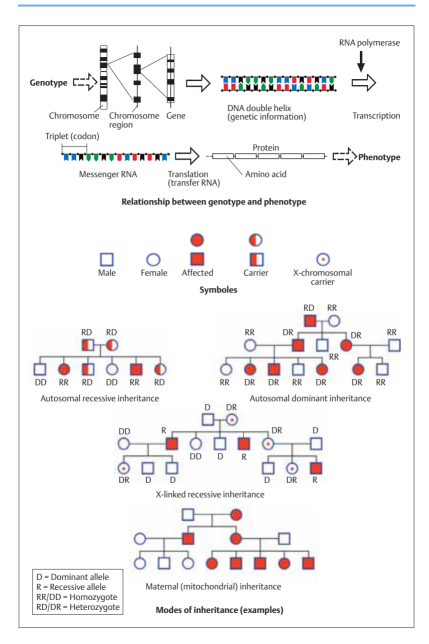
Malformation. A malformation is a structural abnormality of an organ or part of the body in an individual whose body tissues are otherwise normal. Malformations arise during prenatal development because of primary absence or abnormality of the primordial tissue destined to develop into a particular part of the body ("anlage"). *Dysplasia* is malformation due to anomalous organization or function of tissues and tissue components; disorders involving dysplasia include tuberous sclerosis, neurofibromatosis, migration disorders, and various neoplastic diseases.

Developmental anomaly. Disruption of the growth of an organ or body part after normal (primary) primordial development can cause a secondary developmental anomaly. Mechanical influences during development can cause an anomalous position and shape (deformity) of an organ or body part.

■ Infantile Cerebral Palsy (CP) (p. 291)

Infantile cerebral palsy (cerebral movement disorder) is a manifest, but not necessarily unvarying, motor and postural disorder caused by nonprogressive damage to the brain before, during, or after birth. The underlying brain damage is usually of multifactorial origin. *Prenatal* causes include chromosomal defects, infection, hypoxia, or blood group intolerance; *perinatal* causes include hypoxia, cerebral hemorrhage, birth injury, adverse drug effects, and kernicterus; *postnatal* causes include meningoencephalitis, stroke, brain tumor, metabolic disturbances, and trauma.

Symptoms and signs. Paucity of spontaneous movement, abnormal patterns of movement, and delayed development of standing and walking are noted just after birth and as the child develops. Cerebral palsy frequently involves central paresis (hemiparesis, paraparesis, or quadriparesis), spasticity, ataxia, and choreoathetosis (p. 66). There may also be mental retardation, epileptic seizures, behavioral disturbances (restlessness, impulsiveness, lack of concentration, impaired affect control), and impairment of vision, hearing, and speech. The motor disturbances produce deformities of the bones and joints (talipes equinus, contracture, scoliosis, hip dislocation).



Treatment. Physical, occupational, and speech therapy and perception training should be started as soon as possible. Botulinum toxin can be useful in the treatment of spasticity at certain sites (dynamic talipes equinus, leg adductors, arm flexors). *Other measures:* Orthopedic care, seeing and hearing aids, developmental support.

Hydrocephalus

Hydrocephalus is dilatation of the cerebral ventricles (p. 8) due to obstruction of CSF outflow (p. 162). Common etiologies include aqueductal stenosis, Dandy–Walker and Chiari malformations, infection (toxoplasmosis, bacterial ventriculitis), hemorrhage, and obstructing tumors (colloid cyst of the third ventricle, midline tumors).

Symptoms and signs. If the cranial sutures have not yet fused, congenital obstructive hydrocephalus produces an enlarged head (macrocephaly) with a protruding forehead, the result of chronic intracranial hypertension. The head circumference should be measured regularly, as it is a more useful indicator of congenital hydrocephalus than the clinical signs of intracranial hypertension (p. 158), which are often not very pronounced in infants and may be masked by irritability, failure to thrive, crying, and psychomotor developmental delay. These signs include distended veins visible through the patient's thin scalp; bulging of the fontanelles, and vertical gaze palsy (the lower lid covers the open eye to the pupil, the upper lid reveals a portion of the sclera ⇒ "sunsetting"). Signs of intracranial hypertension are the most useful indicators of hydrocephalus once the cranial sutures have fused. A chronic form of hydrocephalus to be differentiated from NPH (p. 160) has been described long-standing overt as ventriculomegaly in adults (LOVA hydrocephalus); symptoms include macrocephaly, headache, lightheadedness, gait disturbances, and bladder dysfunction.

Treatment. Acute hydrocephalus: There is a limited role for medical treatment (e.g., with carbonic anhydrase inhibitors and osmodiuretic agents); neurosurgical treatment is generally needed for CSF drainage (external drainage or surgical shunt) and/or the resection of an obstructing lesion.

Porencephaly

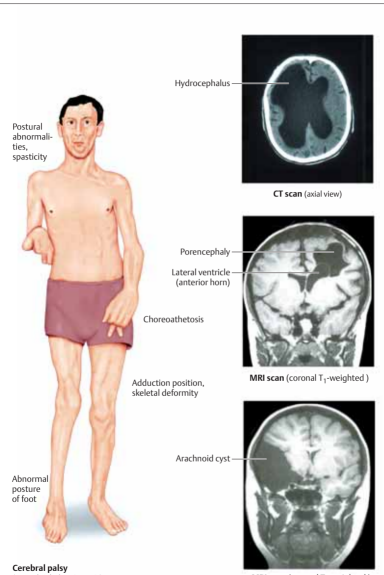
Porencephaly (from Greek *poros*, "opening"), the formation of a cyst or cavity in the brain, is usually due to infarction, hemorrhage, trauma, or infection. Porencephaly in the strict sense of the term involves a communication with the ventricular system. Porencephalic cysts are only rarely associated with intracranial hypertension. Large ones reflect extensive loss of brain tissue; the extreme case is termed *hydranencephaly*. Porencephaly may be asymptomatic or may be associated with focal signs (paresis, epileptic seizures).

Arachnoid Cysts

An arachnoid cyst is a developmental anomaly of the leptomeninges (p.6), usually supratentorial, and located either within the leptomeningeal membranes or between the arachnoid and pia mater. Some arachnoid cysts communicate with the subarachnoid space. Many are asymptomatic, even when large. In rare cases, they can obstruct the CSF pathways (midline or infratentorial arachnoid cysts) or cause new or progressive signs and symptoms because of intracystic hemorrhage, cyst expansion (perhaps by a one-way valve mechanism), or cyst rupture. Symptomatic arachnoid cysts are treated neurosurgically by shunting, fenestration, or excision.

Agenesis of the Corpus Callosum

Hypoplasia or agenesis of the corpus callosum occurs as an isolated finding or in combination with other anomalies (Chiari malformation, heterotopy, chromosomal anomaly, Aicardi syndrome ⇒ infantile spasms, micro-ophthalmia, chorioretinopathy, costovertebral anomalies). Isolated agenesis of the corpus callosum may be asymptomatic and is occasionally found incidentally on CT or MRI scans. Cystic deformities of the septum pellucidum (cavum septi pellucidi, cavum vergae) may obstruct the flow of CSF and cause intracranial hypertension.



(central right hemiparesis)

MRI scan (coronal T₁-weighted)

Anomalies of the Craniocervical Junction

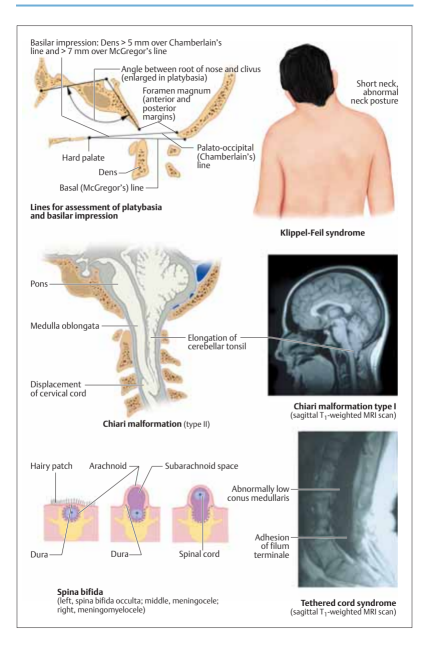
Syndrome	Symptoms and Signs	Causes	Diagnosis/Treatment
Platybasia	Usually asymptomatic	Flattening of the skull base	Plain radiograph ¹ /None
Occipitalization of C1	Usually asymptomatic; possible signs of medullary dysfunction	Synostosis of C1 with the occiput	Plain radiograph, CT, MRI/ Surgical decompression if symptomatic
Basilar impression	Occipitocervical pain; reduced neck flexibility. Long-term: im- pairment of gait, urinary reten- tion, dysarthria, dysphagia, ver- tigo, nausea	Underdevelopment of the occipital bone causing "elevation" of cervical spine ²	Plain radiograph, CT, MRI/ Usually symptomatic; medullary symptoms ⇔ neurosurgical treatment
Klippel–Feil syn- drome ³	Short neck, abnormal head pos- ture, high shoulders, headache, radicular symptoms in arm; possible spinal cord compression	Fused cervical vertebrae	Same as above/ Treatment depends on signs and symptoms

1 Angle between root of nose and clivus > 145°. 2 Congenital (Chiari malformation), acquired (Paget disease, osteomalacia). 3 Additional malformations such as syringomyelia, spina bifida, cleft palate, or syndactyly may be present.

Spinal Dysraphism (Neural Tube Defects)

Syndrome	Symptoms and Signs	Causes	Diagnosis/Treatment
Anencephaly	Absence of cranial vault; cerebral aplasia; normally developed viscerocranium	Nonclosure of anterior por- tion of neural tube	Prenatal ultrasound screen- ing/Termination of preg- nancy
Encephalocele	Protrusion of brain tissue through a midline skull defect ¹	Inhibition malformation (incomplete closure of neural tube)	Measurement of α -feto- protein ² , prenatal ultra- sound screening/Folic acid- vitamin B ₁₂ administration during pregnancy; surgical repair if indicated
Dandy–Walker mal- formation	Hydrocephalus, hypoplasia/ agenesis of vermis; cystic dilata- tion of 4th ventricle; variable degree of facial dysmorphism	Abnormality of embryonal development	CT, MRI/Shunt
Chiari malformation ³	Lower cranial nerve and brain- stem dysfunction (dysphagia, res- piratory dysfunction), head, neck and shoulder pain; abnormal head posture, vertigo, downbeat nystagmus, hydrocephalus (type II)	Abnormality of early embryonal development (weeks 5–6 of gestation)	CT, MRI/Suboccipital decompression; shunt pro- cedure for hydrocephalus; early surgery for myelo- meningocele
Spina bifida4	Spina bifida occulta: Dermal sinus, lumbar hypertrichosis, lum- bosacral fistula, leg pain, gait dis- turbance, foot deformities, blad- der dysfunction (enuresis in children) Other forms: Sensorimotor para- plegia at birth; bladder/bowel dysfunction, foot deformities; hy- drocephalus may occur	Inhibition malformation (incomplete closure of neural tube)	iΩα-Fetoprotein², prenatal ultrasound screening, plain X-ray, CT, MRI/Folic acid administration during pregnancy; surgical treat- ment, physiotherapy, or- thopedic therapy
Tethered cord syndrome	Same as above, with varying severity. Low-lying conus medul- laris, fixed filum terminale	Traction on spinal cord and cauda equina	MRI/Surgery for symptoms and signs reflecting dys- function of the spinal cord and/or cauda equina

1 Meningocele: Only the meninges protrude through the skull defect. Meningoencepholocele: Meninges + brain. Meningoencepholocystocele: meninges + brain + ventricular system. 2 In maternal serum; also in amniotic fluid in open defects. 3 Type is unilateral or bilateral cerebellar tonsillar hemiation with or without caudal displacement of medulla; hydrocephalus, syrin-gomyelia (p. 284); there may be an accompanying anomaly of the skull base. Type II: same as type I + caudal displacement of medulla; hydrocephalus, syrin-gomyelia (p. 284); there may be an accompanying anomaly of the skull base. Type II: same as type I + caudal displacement of medulla; hydrocephalus, syrin-cephalocele. 4 Rachischisis = fissure of vertebral column, incomplete closure of neural tube; spina bifda occulta = incomplete vertebral arch (lamina) with normal position of spinal cord and meningos; meningocele = the arachnoid lies directly under the skin, not covered by the missing dura and bone; myelomeningocele = prolapse of spinal cord (or cauda equina) and arachnoid through the dural and bony defect; diastematomyelia = spilt spinal cord, with two halves separated by connective tissue or a bone spur.



The Phakomatoses

The phakomatoses (neurocutaneous diseases) are a group of congenital diseases in which pathological changes are found in both the central nervous system and the skin. Neurofibromatosis, tuberous sclerosis, and von Hippel–Lindau disease are transmitted in an autosomal dominant inheritance pattern with high penetrance and variable phenotypic expression. These disorders are generally characterized by the formation of benign nodules (hamartoma); malignant tumors (e.g., hamartoblastomas) are rare.

Neurofibromatosis (NF; von Recklinghausen Disease)

The genetic locus for neurofibromatosis type 1 (NF1), the "classical" form of the disease, is on chromosome 17q11.2; that for NF type 2 (NF2) is on 22q12.2.

Symptoms and signs. The characteristic lesions of NF1 are found in the skin (early stage: caféau-lait spots, axillary/inguinal freckling; later stages: neurofibromas/plexiform neurofibromas), eyes (Lisch nodules = whitish hamartomas of the iris, optic glioma), and bone (cysts, pathological fractures, skull defects, scoliosis). There may also be syringomyelia, hydrocephalus, epileptic seizures, precocious puberty, or pheochromocytoma. The hallmark of NF2 is bilateral acoustic neuroma with progressive bilateral hearing loss. Cutaneous manifestations are rare; other nervous system tumors (neurofibroma, meningioma, schwannoma, glioma) are more common. Subcapsular cataract is a typical feature of NF2 in children.

Treatment. Symptomatic tumors are resected.

Tuberous Sclerosis (TSC; Bourneville–Pringle Disease)

The clinical syndrome of tuberous sclerosis is produced by a mutation at either one of two known loci (TSC1: 9q34, TSC2: 16p13.3). TSC1 and TSC2 are clinically identical.

Symptoms and signs. Epileptic seizures (infantile spasms and salaam seizures = West syndrome; focal, generalized) are found in association with *skin changes* (early: hypomelanotic linear spots readily visible under UV light; late signs: adenoma sebaceum, subungual angiofibroma, thick and leathery skin in the lumbar region), *ocular* changes (retinal hamartoma), and tumors (cardiac rhabdomyoma, renal angiomyolipoma, cysts). There may be marked mental retardation and behavioral abnormalities (vocal and motor stereotypy, psychomotor restlessness). CT and MRI reveal periventricular calcification, cortical lesions. and tumors.

Treatment. Symptomatic (anticonvulsants).

Von Hippel–Lindau Disease

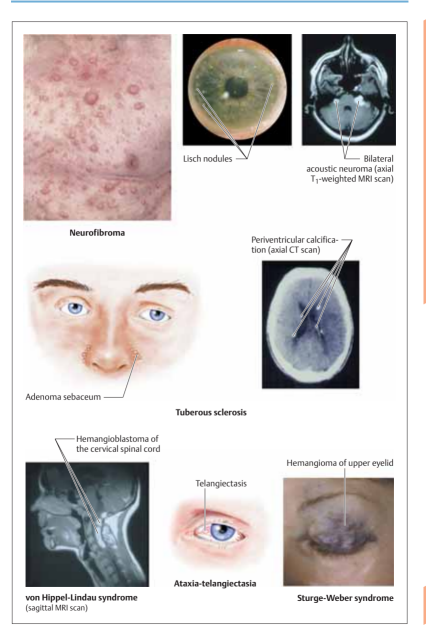
Gene locus. 3p25-p26.

Symptoms and signs. Cystic cerebellar hemangioblastoma causes headache, vertigo, and ataxia, and possibly hydrocephalus by compression of the 4th ventricle. Hemangioma may also occur in the spinal cord. Further lesions are often present in the *eyes* (retinal angiomatosis ⇒ retinal detachment), *kidneys* (cysts, carcinoma), *adrenal glands* (pheochromocytoma), *pancreas* (multiple cysts), and *epididymis* (cystadenoma). **Treatment**. Regular screening of each potentially involved organ system is carried out so that tumors can be resected and vascular complications prevented as early as possible.

Cutaneous Angiomatoses with CNS Involvement

Sturge–Weber disease (encephalofacial angiomatosis). A unilateral or bilateral port wine stain (nevus flammeus) is present at birth and may be either localized (characteristically in the upper eyelid and forehead, in which case involvement of the brain is likely) or widespread (entire head or body). Not all cutaneous hemangiomas are accompanied by cerebral involvement.

Hereditary hemorrhagic telangiectasia (HHT; Osler–Weber–Rendu disease). Known genetic loci: 9q34.1 (HHT1) and 12q11–14 (HHT2). Telangiectases (vascular anomalies) of the skin, mucous membranes, gastrointestinal tract, urogenital tract, and CNS cause recurrent bleeding (nosebleed, gastrointestinal hemorrhage, hematuria, hemoptysis, cerebral hemorrhage, anemia). Arteriovenous shunting in the lung may cause cyanosis and polycythemia.



Throughout the industrialized world, the population is becoming older. It is predicted that the percentage of persons over age 65 will rise further, while that of persons under age 15 will fall.

Aging

Aging is a biological process with a characteristic temporal course. Senescence refers to the physical changes associated with aging. It is still unclear whether human aging is specifically genetically predetermined or, alternatively, reflects cumulative damage incurred over time. The cellular correlates of aging include an increase in spontaneous chromosomal mutations, altered protein conformations, impairment of cell metabolism by the accumulation of free oxygen radicals, a decline of mitochondrial function with an increase in apoptosis (genetically programmed cell death), and diminished activity of regenerative processes. It is not yet known whether these processes are the cause or the effect of aging.

The current average natural lifespan, barring premature death from disease or external causes, is approximately 85 years. The maximum human lifespan (to date, at least) is approximately 120 years. Life expectancy is the average statistically predicted lifespan of a given population at a given point in time. Human life expectancy has risen considerably in the course of history, particularly in the 20th century. The active life expectancy, or the expected time during which the individual can function independently with regard to meals, dressing, personal hygiene, shopping, and finances, is of practical significance. About 35% of persons over 85 are not fully independent, and about 20% need nursing-home care.

Aging and Disease

Aging decreases physiological reserve, i.e., the ability to compensate for the effects of harmful influences, be they endogenous (e.g., diabetes mellitus, heart failure, thyroid dysfunction) or exogenous (e.g., trauma, infection, side effects of medication). Diseases therefore tend to affect the elderly with shorter latency and greater severity. Age-related changes promote the development of diseases such as Alzheimer disease or stroke, as well as accidents such as falls. Brain tumors (p. 254), particularly metastases, glioma, meningioma, acoustic neuroma, and primary cerebral lymphoma, are more common in older patients. Aging is neither a disease nor a cause of disease, but it increases the chance of becoming ill. The physician must distinguish changes due to disease from those of normal aging (see Table 41, p. 382).

Aging and Degenerative Changes

Structural changes in the brain due to aging rather than disease are referred to as involution. Gross changes include diminished brain volume. gyral atrophy, ventriculomegaly, leukoaraiosis (p. 298), parasagittal leptomeningeal fibrosis, and ventricular expansion, while microscopic changes include neuron and axon loss, gliosis, and the presence within neurons of lipofuscin, neuromelanin, granulovacuolar degeneration, microtubular neurofibrillary tangles (NFTs), senile plaques, Lafora bodies, and Lewy bodies. The term abiotrophy refers to the genetically determined. age-dependent occurrence of degenerative changes such as these. Degenerative diseases, on the other hand, are characterized by an abnormal accentuation of these and other morphological changes, producing typical constellations of functional disturbances (disease-specific clinical syndromes). They develop slowly, progressively, sometimes asymmetrically, and with variable intervals of relatively stable disease manifestations. Some are familial.

Alzheimer Disease (AD)

Alzheimer disease (in its sporadic form) is the most common cause of dementia in old age (p. 136). It progresses steadily or stepwise and usually leads to death in 8–10 years (range, 1–25 years). Risk factors for AD include old age, family history of AD, female sex, elevated plasma homocysteine concentration, and the presence of the allele Apo E&A. Nonsteroidal anti-inflammatory drugs appear to lower the risk of AD. Familial AD is rare; it is transmitted in an autosomal dominant inheritance pattern.

Symptoms and Signs

Early stage. Memory impairment develops gradually and almost imperceptibly, barely differing at first from that of benign senile forgetfulness (p. 136). Ultimately, however, the cognitive deficits of AD produce noticeable changes of behavior, e.g., when the patient is working, shopping, running errands, or taking care of finances, or operating such devices as a telephone, stove, television, or computer. The patient may be aware of these deficits and become additionally anxious and depressed because of them. The distinction between AD with depressive features and primary depression with secondary cognitive changes (pseudodementia) has major implications for treatment (Table 42, p. 383).

Intermediate stage. The patient is too confused and disoriented to carry out previous occupational and social activities (p. 132) and needs help and supervision in almost all activities, but may still be able to perform habitual daily routines, carry on a simple conversation, and abide by the basic rules of etiquette. *Aphasia* (e.g., impaired comprehension of speech, word-finding difficulty, cf. p. 126) and *apraxia* (p. 128) are often present. The patient cannot do simple arithmetic or tell time. Visual agnosia is rare at this stage.

Late stage. The increasing cognitive impairment and loss of reasoning and judgment make it impossible for the patients to plan their activities. The patient's aimless wandering, undirected motor activity, and inability to recognize people-even close relatives and friendscomplicate the caregiver's job, along with changes in the patient's circadian rhythm (quiet or apathetic by day, restless by night), impulsive behavior (packing suitcases or running away), delusions, hallucinations, paranoid suspicion of close relatives and friends, aggressive behavior, and neglect of personal hygiene. Patients with advanced AD need help with the simplest activities of daily living (eating, dressing, going to the toilet). They may become incontinent of urine and stool, bedridden, akinetic, and mute. Pathological reflexes (sucking and grasping) can be elicited. Auditory and tactile stimuli may trigger epileptic seizures and myoclonus (for differentiation from

Creutzfeldt–Jakob disease, see p. 252). Death is caused by secondary complications such as pneumonia and heart failure.

Pathogenesis

PET and SPECT studies in early AD reveal bilateral metabolic disturbances in the parietotemporal cortex and decreased neurotransmitter activity in cortical cholinergic fibers (acetylcholine, choline acetyltransferase, nicotinic acetylcholine receptors ⇒ nucleus basalis, medial septal region), serotonergic fibers (raphe nuclei), and noradrenergic fibers (locus coeruleus). There is probably a reduction of cortical glutamatergic activity (excitatory) with a preponderance of GABAergic activity (inhibitory). Neuropathology: Changes such as neuronal death, neuritic (senile) plagues (NPs), and intraneuronal neurofibrillary tangles (NFTs) are seen mainly in the entorhinal cortex, hippocampus, temporal cortex, primary/secondary visual cortex, and nucleus basalis. NPs consist of a central core containing amyloid-Aβ, apolipoprotein E (Apo E), α_1 -antichymotrypsin, synuclein, and other proteins, surrounded by dead neurons, activated glial cells, macrophages, and other inflammatory cells. NFTs consist of paired helical filaments (PHFs) composed of tau proteins, which are normally an important stabilizing component of the microtubular (neurofibrillary) cytoskeleton. Increased phosphorylation of tau proteins leads to the formation of NFTs. Amyloid-Aß (normal function unknown) is formed by proteolysis of the transmembrane amyloid precursor protein (APP), to which neurotrophic and neuroprotective properties have been ascribed. A point mutation in APP on chromosome 21q has been implicated in familial AD; patients over 40 years of age with Down syndrome (trisomy 21) also have neuropathological changes similar to those of AD. Accumulation of amyloid-Aß in arterial walls is the basic abnormality in amyloid angiopathy (p. 178). The gene for Apo E (a lipoprotein involved in cholesterol transport) is found on chromosome 19q and has three alleles, designated £2, £3, and £4; £4 is strongly associated with both sporadic and familial AD. Other familial forms of AD have been traced to mutations of the presenilin-1 gene (PS1 \Rightarrow 14q24.3 \Rightarrow protein S182) and the presenilin-2 gene ($PS2 \Rightarrow 1q31-42 \Rightarrow STM2$ protein). These genes encode cytoplasmic neuronal proteins whose function is as yet unknown.

Treatment

There is no specific treatment for AD. Symptomatic, social, and psychiatric measures and family assistance are the mainstays of treatment. Acetylcholinesterase inhibitors (donepezil, galantamine, rivastigmine, tacrine) or N-methyl-D-aspartate (NMDA) inhibitors (memantine) can improve cognitive function in the early stages of disease. A proposed protective effect of estrogen therapy in postmenopausal women has not been confirmed.

Pick Disease, Frontotemporal Dementia (FTD)

Pick disease causes behavioral changes (e.g., reduced interpersonal distance, apathy, abulia, obsessive-compulsive symptoms, amnestic aphasia, increased appetite) and impairment of semantic memory. CT and MRI mainly reveal asymmetric frontotemporal atrophy. Neuropathology: Neuron loss, gliosis, and variably severe "ballooning" of neurons (Pick cells). There are no neuritic plaques. The disease is familial in 40% of cases: familial Pick disease is transmitted in an autosomal dominant inheritance pattern and is due to a mutation on chromosome 17 (FTDP-17) that causes changes in tau protein (=> FTD with parkinsonian manifestations).

Vascular Dementia (Table 14, p. 367)

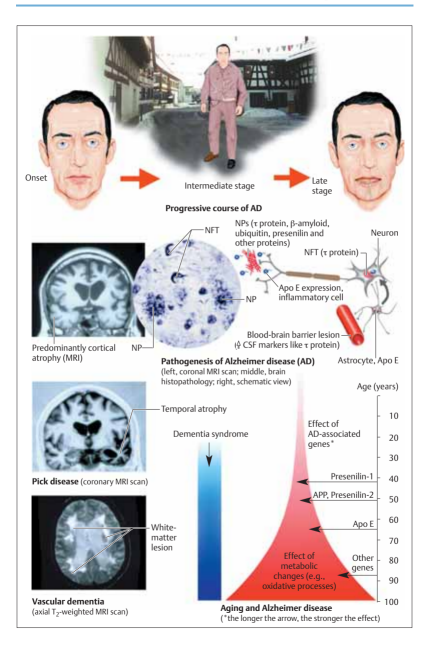
Cerebrovascular disturbances can produce dementia in a variety of ways. The main risk factors for cerebrovascular dementia are old age, arterial hypertension, diabetes mellitus, and generalized atherosclerosis. After AD, cerebrovascular disturbance are the second most common cause of dementia.

Multi-infarct dementia. Multiple small infarcts (lacunes) or large bilateral infarcts may produce any of a variety of focal neurological, behavioral, and cognitive disturbances, depending on their location and extent. These disturbances usually progress in stepwise fashion. CADASIL (p. 172) is a rare cerebrovascular disorder that predisposes to multi-infarct dementia.

Subcortical arteriosclerotic encephalopathy (SAE) is characterized by rarefaction of the white matter (leukoaraiosis) due to microangiopathy. Neuropathology: Histological examination reveals demvelination and reactive gliosis in the white matter, along with changes in the walls of small arteries (hvalinosis, fibrinoid necrosis, hypertrophy). These vascular changes are the cause of chronic ischemic and secondary metabolic damage in areas of white matter supplied by terminal branches. Behavioral changes (attention deficit, loss of cognitive flexibility, abulia, disorientation), gait disturbances, pseudobulbar palsy, urinary incontinence, and other neurological deficits develop slowly and continuously (not in stepwise fashion, as in multiinfarct dementia).

Strategic infarct dementia. Dementia can be produced by a localized infarct in a particular, "strategic" areas of the brain (e.g., limbic system, thalamus, cortical association areas).

Leukoaraiosis is characterized by white-matter lesions (WMLs) that are hypodense on CT and hyperintense on T2-weighted MRI. The extent of WMLs is correlated with the clinical severity of the disease: there may be mild or moderate cognitive impairment (cognitive slowing, memory loss) or severe dementia. WMLs are not always due to a cerebrovascular disturbance and are present in a variety of conditions other than chronic arterial hypertension (e.g., AD, multiple sclerosis. PML, Creutzfeldt-Jakob disease, ADEM, trauma, radiation therapy, chemotherapy, vitamin B₁₂ deficiency, hypoxic-ischemic encephalopathy, CADASIL, central amyloid angiopathy).



Huntington Disease (HD)

The first symptoms of HD typically appear between the ages of 35 and 45 years. HD appearing before age 20 (the Westphal variant of HD) is characterized by akinesia, bradykinesia, epileptic seizures, action tremor, and myoclonus. Onset before age 10 or after age 70 is rare. HD patients require total nursing care 10–15 years after the onset of this inexorably progressive, and ultimately fatal degenerative disease.

Symptoms and Signs

Early stage. In cases of earlier onset, akinesia and cognitive impairment tend to be more prominent than choreiform movements, while, in cases of later onset, the reverse is true, Behavioral changes such as depression, suicidal tendencies, paranoia, querulousness, irritability, impulsiveness, emotional outbursts, aggressive behavior, poor hygiene, loss of initiative, and inappropriate sexual behavior impair familial and social relationships and may even lead to criminal charges. Other changes include cognitive slowing, diminished tolerance for stress, and impairment of memory and concentration. The patient becomes obviously unable to perform his or her usual tasks at work or at home. Chorea (p. 66; Table 43, p. 383) may initially be misdiagnosed as "nervous" agitation or fidgeting. Even severe chorea disappears during sleep. Chorea may be accompanied by akinesia, dystonia, and decreased voluntary motor control. The patient's gait is impaired by poor balance and loss of postural motor control. Oculomotor disturbances are also common.

Intermediate stage. Progressive dementia (p. 136) is accompanied by loss of drive, generalized choreiform, dystonic, and bradykinetic movements, and frequent falls.

Late stage. Many patients become cachectic, with muscular atrophy (⇔ interosseous muscles of hands) and weight loss despite an adequate caloric intake. Chorea is largely replaced by akinesia in late HD. General motor control is greatly impaired. Urinary incontinence is not uncommon. These patients need full nursing care.

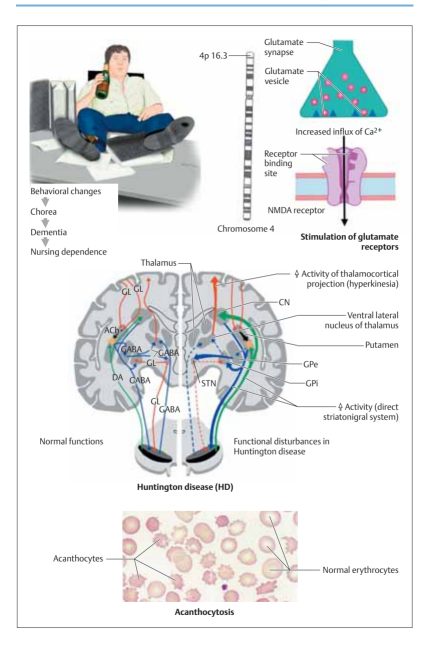
■ Pathogenesis (for abbreviations, see p. 211)

HD is characterized by generalized cerebral atrophy, especially of the dorsal striatum

(p. 210), and, neurochemically, by a marked deficiency of GABA and of glutamate decarboxylase (an enzyme involved in GABA synthesis). HD is transmitted in an autosomal dominant pattern with complete penetrance, that is, all persons bearing the gene eventually develop the disease. The gene for HD (IT15) is on the end of the short arm of chromosome 4 (4p16.3), which contains CAG repeats (the trinucleotide sequence CAG codes for glutamine; cf. p. 288). Healthy subjects have 11-34 CAG repeats at this locus, while persons with HD have more than 40. Paternal inheritance is associated with anticipation (increasingly early onset in subsequent generations), but maternal inheritance is not. The gene product is referred to as huntingtin. The pathophysiological mechanism of HD remains obscure; increased glutamatergic transmission at NMDA receptors is thought to produce neurodegenerative changes (excitotoxicity). There is now a direct gene test that can be performed on a peripheral blood sample to detect HD before the onset of symptoms. It may only be performed with the informed consent of a patient above the legal age of majority, after the potential social and psychiatric implications of a positive result have been explained.

Neuroacanthocytosis

Acanthocytes (crenated erythrocytes with thorny processes) make up more than 3% of all erythrocytes in fresh blood smears obtained from patients with the following neurological diseases: *abetalipoproteinemia* (Bassen-Kornzweig syndrome, p. 280, 307; © cholesterol and triglycerides), *McLeod syndrome* (X-linked recessive myopathy; absence of Kell precursor protein), and *neuroacanthocytosis* (normal lipoprotein concentrations). The mode of inheritance of neuroacanthocytosis is unknown. Its major features, orofacial dyskinesia (tonguebiting and lip-biting) and chorea, usually appear between the ages of 20 and 30 years.



Atypical Parkinsonian Syndromes

Roughly 80% of patients with parkinsonism suffer from *idiopathic Parkinson disease* (p. 206 ff), while the rest have either *symptomatic parkinsonism* (Table 44, p. 383) or *atypical parkinsonism* (AP) due to one of the neurodegenerative "Parkinson-plus" syndromes. The more common AP syndromes are multiple system atrophy, progressive supranuclear palsy, and corticobasal degeneration. An unequivocal differentiation of these disorders from one another, and from idiopathic Parkinson disease, may not be possible at the onset of symptoms, or even later in the course of the disease.

Multiple System Atrophy (MSA)

MSA is a gradually progressive, sporadic, nonfamilial disease of adults marked by autonomic and cerebellar dysfunction of variable severity, accompanied by parkinsonian manifestations that respond poorly to levodopa. The term MSA covers the earlier-described disorders olivopontocerebellar atrophy (OPCA), idiopathic orthostatic hypotension (IOH), Shy-Drager syndrome, and striatonigral degeneration (SND). The onset of symptoms is usually between the ages of 45 and 70. Autonomic dysfunction is manifested by urinary incontinence (p. 156; abnormal EMG of urethral/anal sphincter ⇒ increased polyphasic rate) and orthostatic hypotension (p. 148), and sometimes by hypertension while the patient is down. Cerebellar dysfunction lving is manifested by gait ataxia, frequent falls, dysarthria, and oculomotor disturbances. The parkinsonian manifestations include akinesia, rigidity, postural instability (p. 206), and frequent falling, but there is no resting tremor.

Progressive Supranuclear Palsy (PSP)

PSP (Steele–Richardson–Olszewski syndrome) is a progressive, nonfamilial disease that usually appears around the age of 40. Its major features are unsteady gait, postural instability, frequent falls (often backward), axial dystonia, rigidity, akinesia, behavioral changes (bradyphrenia, irritability, social withdrawal, abnormal fatigability, uncontrollable laughing and crying), and pseudobulbar palsy (p. 166). Paralysis of voluntary conjugate upward gaze may be present at onset, but paralysis of downward gaze is more common. Passive (reflex) vertical eye movements (doll's eye sign) are still present, i.e., the vestibulo-ocular reflex remains intact (p. 26). The fully developed clinical picture of PSP (abnormally erect posture, retrocollis, wide-open eyes, elevated forehead muscles, dysarthria, dysphagia, and frontal brain syndrome, p. 122) is practically pathognomonic, but its manifestations in the early phase can be very difficult to distinguish from those of other neurodegenerative diseases. The symptoms and signs of PSP respond poorly to levodopa, if at all.

Corticobasal Degeneration (CBD)

CBD is a progressive neurodegenerative disease that most often occurs in adults over the age of 60. Its major features are akinesia, rigidity, limb apraxia (p. 128), and cortical sensory deficits (astereognosis, graphanesthesia). There is a loss of motor control, so that the patient's hands (limbs) appear to move spontaneously without the patient's guiding them (*alien hand/limb phenomenon*). Gait instability is an early sign and is later accompanied by dysarthria, dysphagia, myoclonus, dystonic arm posture (wrist and elbow flexion, shoulder adduction), action/postural tremor, supranuclear oculomotor disturbances (p. 86), blepharospasm, and cognitive impairment. Levodopa is unhelpful.

Dementia with Lewy Bodies (DLB), Diffuse Lewy Body Disease

DLB is characterized at first by an akinetic-rigid parkinsonian syndrome (p. 208), which is later accompanied by fluctuating behavioral changes (attention deficit, disorientation, impairment of consciousness, visual hallucinations) and frequent falling for no apparent reason. Patients are hypersensitive to neuroleptics and benzodiazepines. An abundance of Lewy bodies (p. 211) can be observed, particularly in cortical neurons.



Autonomic dysfunction (orthostatic hypotension, urinary incontinence, impotence, anhidrosis)

Asymmetrical parkinsonism, akinesia/rigidity, early postural/gait instability (=> falls)



Ataxia, dysarthria, dysphonia

Multiple system atrophy



Doll's eyes phenomenon on vertical head movement

Vertical gaze palsy (progressive supranuclear palsy)

Limb apraxia, dystonic arm position

Corticobasal degeneration

303

Myoclonus

Motor Neuron Diseases

These diseases involve a degeneration of the cerebral and/or spinal motor neurons (p. 44). They present with a wide variety of neurological syndromes of varying temporal course.

Upper Motor Neuron Diseases

(p. 46; Table 45, p. 384)

Hereditary. Familial spastic spinal paralysis (p. 286), adrenomyeloneuropathy, spinocerebellar ataxia type 3 (p. 280).

Acquired. Lathyrism (central spastic paraparesis due to a neurotoxin in the pulse *Lathyrus sativus* (grass pea), a dietary staple in certain poor districts in India); konzo (= cassavaism, a toxic reaction to flour made of insufficiently processed cassava, seen in certain parts of Africa); tropical spastic paraparesis (HTLV1-associated myelopathy = HAM).

Lower Motor Neuron Diseases

(p. 50; Table 46, p. 385)

Most cases of spinal muscular atrophy are hereditary. Their clinical features vary according to the age of onset. Acquired forms are rare.

Diseases Affecting Both the Upper and the Lower Motor Neuron

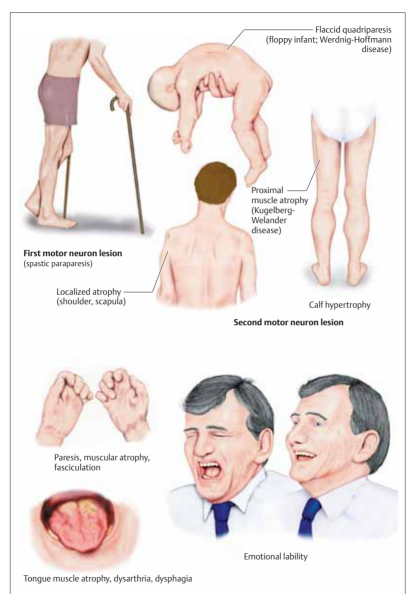
Hereditary. Amyotrophic lateral sclerosis (ALS) is familial in 5-10% of cases. Familial ALS with onset in childhood and adolescence (juvenile ALS) is transmitted either as an autosomal recessive trait (ALS2:2q33;ALS5:15q15.1-21.1) or as an autosomal dominant trait (9q34 linkage). Adult-onset familial ALS is transmitted as an autosomal dominant trait (ALS1:21q22.1; ALS3:18q21) associated with a mutation of the gene for superoxide dismutase 1 (SOD1). SOD1 plays a role in converting cytotoxic oxygen radicals to hydrogen peroxide. It remains unknown how the SOD1 defect causes motor neuron disease. Autosomal dominant inheritance has also been found for ALS plus frontotemporal dementia (9q21-22). ALS together with parkinsonism and dementia occurs among the Chamorro people of Guam.

Acquired. Sporadic ALS usually becomes apparent between the ages 50 and 70 (Table 47, p. 386). The presentation is typically with asymmetric weakness of the limbs, either proximal

(difficulty raising the arms or standing up from a sitting position) or distal (frequent falls; difficulty grasping, turning a key in a lock), or else with bulbar dysfunction (dysarthria). These deficits are often accompanied by leg cramps and continuous, marked fasciculation in the proximal limb muscles. As the disease progresses, weakness, muscular atrophy, dysphagia, and dysarthria become increasingly severe. Respiratory weakness leads to respiratory insufficiency. Spasticity, hyperreflexia, pseudobulbar palsy, emotional lability, and Babinski reflex (inconsistent) are caused by dysfunction of the first motor neuron; muscular atrophy and fasciculation are caused by dysfunction of the second motor neuron; and dysarthria, dysphagia, and weakness are caused by both. About 10% of patients have paresthesiae, and some have pain in later stages of the disease. Bladder, rectal, and sexual dysfunction, impairment of sweating, and bed sores are not part of the clinical picture of ALS. The disease progresses rapidly and usually causes death in 3-5 years.

Treatment

There is currently no effective primary treatment for motor neuron diseases. Treatment can be provided for the palliation of various disease manifestations, e.g., dysarthria (speech therapy, communication aids), dysphagia (swallowing training, percutaneous endoscopic gastrostomy, surgery), and drooling (medication to decrease salivary flow). Antispasmodic agents can be used to treat spasticity and muscle spasms, and psychiatric medications to treat emotional lability. Physical and occupational therapy are provided, including breathing exercises, contracture prophylaxis, and measures to increase mobility. Further measures include orthoses, breathing training (aspiration prophylaxis, secretolysis, ventilator for home use, tracheotomy), and psychosocial support. Riluzole (a glutamate antagonist) has been found to prolong survival in ALS.



Lesion of both first and second motor neurons

Encephalopathies

The term *encephalopathy* refers to a focal or generalized disturbance of brain function of noninfectious origin. Depending on their etiology, encephalopathies may be reversible, persistent, or progressive. Their clinical manifestations are diverse, depending on the particular functional system(s) of the brain that they affect.

Hereditary Metabolic Encephalopathies

These disorders frequently cause severe cognitive impairment. Most of them have an autosomal recessive inheritance pattern; a few are X-linked recessive. The underlying primary enzyme defect (enzymopathy) may be a monogenic, polygenic, or mitochondrial genetic trait, or a multifactorial disorder (see p. 288). All hereditary metabolic encephalopathies are characterized by chronic progression, recurrent impairment of consciousness, spasticity, cerebellar ataxia, extrapyramidal syndromes, and psychomotor developmental delay. The following tables contain a partial listing of hereditary metabolic encephalopathies (Lyon et al., 1996); for such disorders affecting neonates and infants, see p. 386f. Some of the diseases listed may appear earlier or later than the typical age of onset indicated.

Metabolic Encephalopathies of Infancy (up to age 2 years)

Syndrome	Defect/Enzyme Defect	Symptoms and Signs
Phenylketonuria	Phenylalanine hydroxylase deficiency	Psychomotor retardation, hyperactivity, movement disorders, stereotypic move- ments
Hartnup disease	Impaired renal/intestinal trans- port of neutral amino acids	Reddish, scaly changes of exposed skin, emotional lability, episodic cerebellar ataxia
Gaucher disease (type III, sub- acute neuropathic form)	See p. 387	Generalized seizures, ataxia, myoclonus, progressive mental decline, supranuclear oculomotor disturbances, splenomegaly
Niemann-Pick disease (type C)	Exact defect not known	Mental retardation, seizures, ataxia, dys- arthria, vertical gaze palsy
Metachromatic leukodystro- phy	Arylsulfatase A deficiency	Progressive gait impairment, spasticity, progressive dementia, dysarthria, blind- ness, cerebellar ataxia, polyneuropathy
Leigh disease ¹	No consistent defect ²	Respiratory disturbances, gaze palsy, ataxia, decreased muscle tone, retinitis pigmentosa, seizures

1 Subacute necrotizing encephalomyelopathy. **2** Known defects include mitochondrial respiratory chain defects (complexes IV and V) and protein synthesis defects. The clinical features are heterogeneous. MRI scans show multiple, bilaterally symmetric lesions with sparing of the mamillary bodies. CSF lactate concentration increased. Muscle biopsy reveals no ragged red fibers.

Syndrome	Defect/Enzyme Defect	Symptoms and Signs
Abetalipoproteinemia	See pp. 280, 300	Gait impairment, ataxia, dysarthria, poly- neuropathy, night blindness
Progressive myoclonus epilepsy ¹ with Lafora bodies ²	Lysosomes	Epileptic seizures, myoclonus, dementia, cerebel- lar ataxia, epileptic visual phenomena
Wilson disease (dystonic type)	Copper transport protein ³	Dysfunction/cirrhosis of liver, behavioral changes, facio-oropharyngeal rigidity (dysarthria, dyspha- gia), parkinsonism, tremor, dystonia, Kayser- Fleischer ring (p. 309)
Neuronal ceroid lipofusci- nosis (Spielmeyer–Vogt syndrome)	Storage of lipid pigment in lysosomes	Visual impairment, dysarthria, dementia, epilep- tic seizures, myoclonus, parkinsonism
Panthothenate kinase- associated degeneration ⁸	Accumulation of iron pig- ment in substantia nigra and globus pallidus ⁴	Gait impairment, dystonia, dysarthria, behavioral changes, dementia, retinal depigmentation
Adrenoleukodystrophy ⁵	Peroxisomes (p. 386)	Behavioral changes/dementia, gait impairment, cortical blindness, spastic quadriparesis, deafness, primary adrenocortical insufficiency
Homocystinuria ⁶	Cystathionine β -synthase ⁷	Dementia/behavioral changes, osteoporosis, ectopia lentis
Fabry disease ⁶	α-Galactosidase A (ûglycosphingolipids)	Attacks of pain in digits and abdomen; diffuse angiokeratomas; cataract
Mitochondrial syndromes	See p. 402	See p. 402

■ Metabolic Encephalopathies of Childhood and Adolescence (ages 3–18 years)

1 Other forms (p. 68) include Unverricht-Lundborg syndrome, myoclonus epilepsy with ragged red fibers (MERFF, p. 402), late forms of other lysosome defects (e. g., sialidosis type I, GM₂-gangliosidosis). 2 Cytoplasmic inclusion bodies containing glycoprotein mucopolysaccharides in the brain, muscles, skin, and liver (also called Lafora disease). 3 Autosomal recessive trait, mutation at 13q14.3; U serum ceruloplasmin, ¹D hepatic copper, free serum copper, and urinary copper levels, U rate of incorporation of ⁶⁴Cu in ceruloplasmin, MRI signal changes (striatum, dentate nucleus, thalamus). 4 MRI shows bilaterally symmetric hypointensity of globus pallidus with central zone of hyperintensity ("tiger eye" sign). 5 Adrenomyeloneuropathy, p. 384. 6 Increased risk of stroke. 7 Most common form. 8 Formerly called Hallervorden-Spatz disease.

Metabolic Encephalopathies of Adulthood

Syndrome	Defect/Enzyme Defect	Symptoms and Signs
Metachromatic leuko- dystrophy	Arylsulfatase-A deficiency	Behavioral changes, gait impairment, dementia
Krabbe disease	See p. 387	Gait impairment, spastic quadriparesis, poly- neuropathy, optic nerve atrophy
Adrenoleukodystrophy	Peroxisomes	Adult form extremely rare
Neuronal ceroid lipofusci- nosis (Kufs disease)	Storage of lipid pigment in lysosomes	Type A: Epilepsy, myoclonus, dementia, ataxia Type B: Behavioral changes/dementia, facial dys- kinesia, movement disorders
GM ₁ gangliosidosis	See p. 387	Progressive dysarthria and dystonia
GM ₂ gangliosidosis ¹	See p. 387	Chronic progression (p. 387)
Wilson disease (pseudo- sclerotic type) ²	See above	Postural/intention tremor (beginning in one arm), behavioral changes, dysarthria, dysphagia, mask- like facies, parkinsonism
Gaucher disease type 3	See p. 387	Supranuclear ophthalmoplegia, epileptic seizures, myoclonus, splenomegaly
Niemann–Pick disease (type C)	See p. 387	Cerebellar ataxia, intention tremor, dysarthria, supranuclear vertical gaze palsy
Mitochondrial syndromes	See p. 402	See p. 402

1 Tay-Sachs disease. 2 Westphal-Strümpell disease.

Acquired Metabolic Encephalopathies

Hypoxic-ischemic encephalopathy. An acute lack of oxygen (Pao₂ <40 mmHg), severe hypotension (<70 mmHg systolic), or a combination of the two causes loss of consciousness within minutes. The most important causes of hypoxic and ischemic states are an inadequate pumping function of the heart (as in myocardial infarction, shock, and cardiac arrhythmia), suffocation, carbon monoxide poisoning, respiratory muscle paralysis (as in spinal trauma, Guillain-Barré syndrome, and myasthenia), and inadequate ventilation (as in opiate intoxication). Permanent damage usually does not occur if the partial pressure of oxygen and the blood pressure can be brought back to normal in 3-5 minutes. Longer periods of hypoxia and ischemia are rarely tolerated (except under conditions of hypothermia or barbiturate intoxication); brain damage usually ensues, and may be permanent. Persistent coma (p. 118) with absent brain stem reflexes (pp. 26, 118) once the circulation is restored indicates a poor prognosis; the probable outcome is then a persistent vegetative state or death (p. 120). Patients who regain consciousness may develop various postanoxic syndromes, e.g., dementia, visual agnosia, parkinsonism with personality changes. choreoathetosis, cerebellar ataxia, intention or action myoclonus (Lance-Adams syndrome), and Korsakoff syndrome. Delayed postanoxic syndrome occurs 1-4 weeks after the initial recovery from anoxia and is characterized by behavioral changes (apathy, confusion, restlessness) that may either regress or worsen, perhaps to coma. These changes may be accompanied by gait impairment and parkinsonism. Hypercapnia (î) PaCO₂) due to chronic hypoventilation (as in emphysema, fibrosing alveolitis, or central hypoventilation) causes headache, behavioral disturbances, impairment of consciousness (p. 116 ff), asterixis (p. 68), fasciculations, and bilateral papilledema.

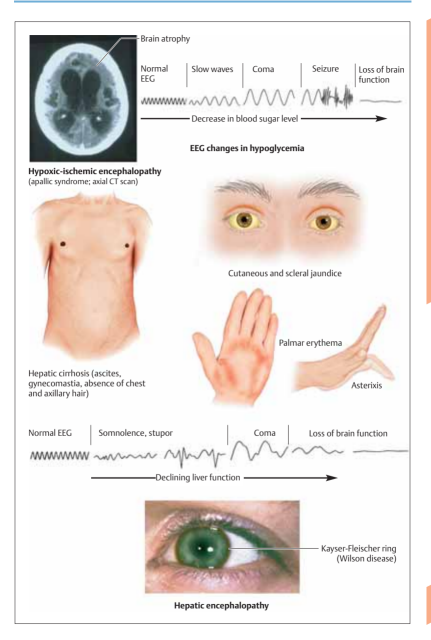
Hypoglycemia. If the blood glucose concentration *acutely* falls below 40 mg/dl, behavioral changes occur (restlessness, hunger, sweating, anxiety, confusion). Any further decrease leads to unconsciousness (grand mal seizure, dilated pupils, pale skin, shallow breathing, bradycardia, decreased muscle tone). Glucose must be given intravenously to prevent severe brain damage. *Subacute* hypoglycemia produces slowed thinking, attention deficits, and hypothermia. *Chronic* hypoglycemia produces behavioral changes and ataxia (p. 324); it is rarely seen (e.g., in pancreatic islet cell tumors).

Hyperglycemia (p. 324). *Diabetic ketoacidosis* is characterized by dehydration, headache, fatigue, abdominal pain, Kussmaul respiration (deep, rhythmic breathing at a normal or increased rate). Blood glucose > 350 mg/dl (\emptyset pH, \emptyset p_{CO₂}, \emptyset HCO₃⁻). In *hyperosmolar non-ketotic hyperglycemia*, the blood glucose con centration is > 600 mg/dl and there is little or no ketoacidosis. The persons at greatest risk are elderly patients being treated with corticosteroids and/or hyperosmolar agents to reduce edema around a brain tumor.

Hepatic/portosystemic encephalopathy occurs by an unknown pathogenetic mechanism in patients with severe liver failure (hepatic encephalopathy) and/or intrahepatic or extrahepatic venous shunts (portosystemic encephalopathy). Venous shunts can develop spontaneously (e.g., hepatic cirrhosis) or be created surgically (portocaval anastomosis, transjugular intrahepatic stent). Clinical features (see Table 50, p. 387): Behavioral changes, variable neurological signs (increased or decreased reflexes, Babinski reflex, rigidity, decreased muscle tone, asterixis, dysarthrophonia, tremor, hepatic coma), and EEG changes (generalized symmetric delta/triphasic waves). The diagnosis is based on the clinical findings, the exclusion of other causes of encephalopathy (such as intoxication, sepsis, meningoencephalitis, and electrolyte disorders), and an elevated arterial serum ammonia concentration.

Repeated episodes of hepatic coma may lead to *chronic encephalopathy* (head tremor, asterixis, choreoathetosis, ataxia, behavioral changes); this can be prevented by timely liver transplantation.

Encephalopathies



Disorders of fluid and electrolyte balance. The regulation of water balance (osmoregulation) is reflected in the serum sodium concentration, [Na⁺]. The hypothalamus, which contains osmoreceptors (p. 142), controls thirst and the secretion of ADH; these in turn determine fluid intake and urine osmolality. Sodium salts account for more than 95% of the plasma osmolality (moles of osmotically active particles per kg of water). Hyperhydration causes a decrease in plasma osmolality (U [Na+]). The ensuing inhibition of thirst and of ADH secretion leads to a reduction of oral fluid intake and to the production of dilute urine (U urinary [Na⁺]), restoring the normally hydrated state. Dehydration induces the opposite changes, again resulting in restoration of the normally hydrated state.

The regulation of sodium balance (volume regulation) maintains adequate tissue perfusion (p. 148). Volume receptors in the carotid sinus and atria of the heart are the afferent arm of the reflex pathway controlling renal sodium excretion, whose efferent arms are the sympathetic system, the renin–angiotensin–aldosterone system (RAAS), and natriuretic peptides. *Hypovolemia and hypervolemia* usually involve combined abnormalities of water and sodium balance.

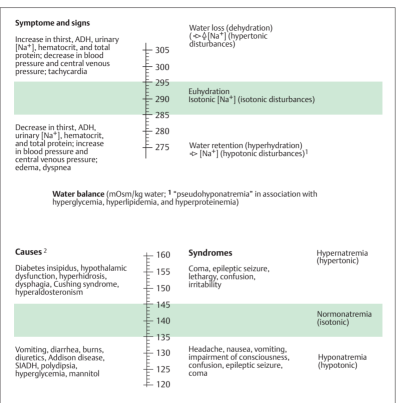
In neurological disorders (head trauma, meningoencephalitis, brain tumor, subarachnoid hemorrhage, acute porphyria), the syndrome of inappropriate ADH secretion (SIADH) is characterized by water retention (volume expansion), abnormally concentrated urine, and hyponatremia. The more rapidly hyponatremia develops, the more severe its clinical signs (e. g., confusion, seizures, impairment of consciousness). SIADH is to be distinguished from central salt-wasting syndrome, which is characterized by hypovolemia and dehydration.

Too rapid correction of hyponatremia causes most cases of *central pontine myelinolysis* (other causes are serum hyperosmolality and malnutrition). In this syndrome (p. 315), a patient with a major systemic illness (e.g., postoperative state, alcoholism) develops quadriplegia, pseudobulbar palsy, and locked-in syndrome (p. 120) over the course of a few days. A less severe form of central pontine myelinolysis is characterized by confusion, dysarthria, and gaze palsies. **Calcium/magnesium.** Hypercalcemia causes nonspecific symptoms along with apathy, progressive weakness, and impairment of consciousness (or even coma). Hypocalcemia is characterized by increased neuromuscular excitability (muscle spasms, laryngospasm, tetany, Chvostek's and Trousseau's signs), irritability, hallucinations, depression, and epileptic seizures. Hypomagnesemia has similar clinical features.

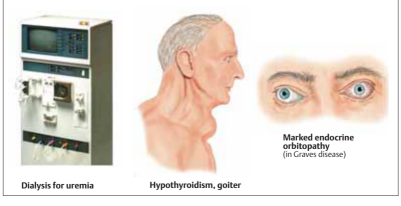
Uremic encephalopathy arises in patients with renal failure and is characterized by behavioral changes (apathy, cognitive impairment, attention deficit, confusion, hallucinations), headache, dysarthria, and hyperkinesia (myoclonus, choreoathetosis, tremor, asterixis). Severe uremia can produce coma. The differential diagnosis of uremic encephalopathy includes cerebral complications of the primary disease, such as intracranial hemorrhage, drug intoxication because of impaired catabolism, and hypertensive encephalopathy. A similar neurological syndrome can arise during or after hemodialysis or peritoneal dialysis (dysequilibrium syndrome). Dialysis encephalopathy (dialysis dementia; now rare) is probably due to aluminum poisoning as a complication of chronic hemodialysis. Its manifestations include dysarthria with stuttering and stammering, myoclonus, epileptic seizures, and behavioral changes (p. 122 ff).

Endocrine encephalopathy is characterized by agitation with hallucinations and delirium, anxiety, apathy, depression or euphoria, irritability, insomnia, impairment of memory and concentration, psychomotor slowing, and impairment of consciousness. It may be produced (in varying degrees of severity) by Cushing disease, high-dose corticosteroid therapy, Addison disease, hyperthyroidism or hypothyroidism, and hyperparathyroidism or hypoparathyroidism.

Encephalopathies



Sodium balance (mmol/L;² examples, some with combined deficits)



Encephalopathies

Encephalopathy due to sepsis, multiple organ failure, or burns may arise within a few hours, manifesting itself as impaired concentration, disorientation, confusion, and psychomotor agitation in addition to the already severe systemic disturbances. In severe cases, there may be delirium, stupor or coma. Focal neurological signs are absent; meningismus may be present, and CSF studies do not show signs of meningoencephalitis. There are nonspecific EEG changes (generalized delta and theta wave activity). The pathogenesis of these syndromes is unclear. Their prognosis is poor if the underlying disease does not respond rapidly to treatment.

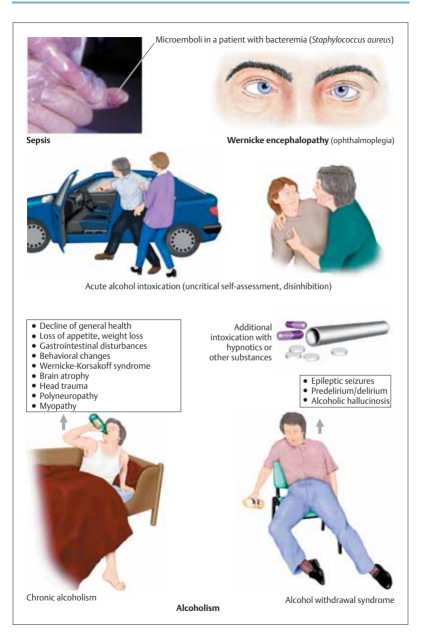
Paraneoplastic encephalopathy occurs as a complication of neoplasms outside the central nervous system. It can only be diagnosed after the exclusion of local tumor invasion or metastasis, complications of tumor treatment, or other complications of the primary disease. For paraneoplastic encephalopathy, see Table 51, p. 388; for paraneoplastic disorders affecting the PNS, neuromuscular junction, and muscle, see p. 406.

Wernicke-Korsakoff syndrome. Wernicke encephalopathy is characterized by gaze-evoked nystagmus or dissociated nystagmus, ophthalmoplegia (abducens palsy, conjugate gaze palsy or, rarely, miosis), postural and gait ataxia, and impairment of consciousness (p. 116; apathy, indifference, somnolence). Korsakoff syndrome is characterized by confabulatory amnesia (p. 134), disorientation, and decreased cognitive flexibility. Most patients have a combination of these two syndromes, which is then called Wernicke-Korsakoff syndrome. Polyneuropathy, autonomic dysfunction (orthostatic hypotension, tachycardia, exercise dyspnea), and anosmia may also be present. These syndromes are caused by a deficiency of thiamin (vitamin B₁) due to alcoholism or malnutrition (malignant tumors, gastroenterologic disease, thiamin-free parenteral nutrition). This, in turn, causes dysfunction of thiamin-dependent enzymes (increase in transketolase, pyruvate decarboxylase, a-ketoglutarate dehydrogenase, and serum pyruvate and lactate: decrease in transketolase activity in erythrocytes). MRI reveals lesions in paraventricular areas (thalamus, hypothalamus, mamillary bodies) and periaqueductal areas (mid brain, motor nucleus of X, vestibular nuclei, superior cerebellar vermis). *Treatment:* Immediate intravenous infusion of thiamin (50– 100 mg) in glucose solution. *Note:* glucose infusion without thiamin in a patient with latent or unrecognized thiamin deficiency may provoke or exacerbate Wernicke encephalopathy.

Encephalopathies Due to Substance Abuse

Alcohol. Acute alcohol intoxication (drunkenness, inebriation) may be mild (blood alcohol 0.1-1.5 \Rightarrow dysarthria, incoordination, disinhibition, increased self-confidence, uncritical selfassessment), moderate (blood alcohol 1.5-2.5 ‰ ⇒ ataxia, nystagmus, explosive reactions, aggressiveness, euphoria, suggestibility), or severe (blood alcohol >2.5% \Rightarrow loss of judgment, severe ataxia, impairment of consciousness, autonomic symptoms such as hypothermia, hypotension, or respiratory arrest). Concomitant intoxication with other substances (sedatives, hypnotics, illicit drugs) is not uncommon. The possibility of a traumatic brain injury (subdural or epidural hematoma, intracerebral hemorrhage) must also be considered. Pathological intoxication after the intake of relatively small quantities of alcohol is a rare disorder characterized by intense outbursts of emotion and destructive behavior, followed by deep sleep. The patient has no memory of these events.

Alcohol withdrawal syndrome. Reduction of alcohol intake or total abstinence from alcohol after chronic alcohol abuse causes acute autonomic disturbances (sweating, tachycardia, insomnia, nausea, vomiting), tremor, impairment of concentration, and behavioral changes. This initial stage of predelirium is followed by a stage of delirium (delirium tremens), in which all of the disturbances listed worsen and are accompanied by visual hallucinations. Epileptic seizures may occur. The course of delirium tremens can be complicated by systemic diseases that are themselves complications of alcoholism (hepatic and pancreatic disease, pneumonia, sepsis, electrolyte imbalances). Auditory alcoholic hallucinosis without autonomic symptoms or disorientation is an unusual form of alcohol withdrawal syndrome.



Encephalopathies

Late complications of alcoholism. Various disorders are associated with chronic alcohol abuse, though alcohol abuse may not be their only causative factor. *Brain atrophy* is often seen in CT or MRI scans and seems to be reversible by abstinence. In *alcoholic dementia*, brain atrophy is accompanied by cognitive impairment; most cases are probably due to Wernicke–Korsakoff syndrome (p. 312). *Cerebellar atrophy* predominantly affects the anterosuperior vermis (\Rightarrow postural and gait ataxia). *Central pontine myelinoly-* sis (p. 310) and tobacco-alcohol amblyopia (bilateral impairment of visual acuity and visual defects, probably due to a combined deficiency of vitamins B_1 , B_6 , and B_{12}) are other late complications of alcoholism. Fetal alcohol syndrome (congenital malformations, hyperactivity, attention deficit, impaired fine motor control) is seen in the children of alcoholic mothers.

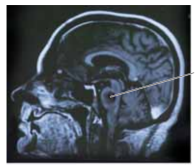
Substance abuse. Neurological signs of substance abuse are described in the table below.

Substance	Pupils	Motor Dysfunction	Reflexes ³	Behavior/Consciousness
Cocaine ^{1,2}	Dilated	Chorea, tremor, dys- tonia, myoclonus, brux- ism	Û	Anxiety, agitation, insomnia, psy- chosis/hypervigilance ⇔ lethargy, coma
Am- phetamines ^{1,2}	Dilated	Chorea, bruxism, muscle spasms, tremor	Û	Euphoria, hyperactivity, dys- phoria, hallucinations, confusion/ hypervigilance
MDMA ^{1,2,4}	Dilated	Tremor, rigidity	Û	Anxiety, hyperactivity, psychosis/ coma ⁵
Opiates ^{1,6}	Pinpoint	Hypokinesia, parkin- sonism	Û	Euphoria/somnolence ⇔ coma, respiratory depression
LSD ⁷	Dilated, slug- gish	Tremor	Û	Euphoria, panic, depression, hal- lucinations, illusions
Phencyclidine (PCP)	Miotic; nystag- mus	Ataxia, tremor, increased muscle tone	Û	Euphoria, dysphoria, psychosis, aggressiveness, hallucinations/ coma (rare)

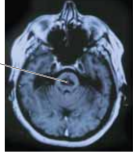
1 Epileptic seizures may occur. 2 Cerebral infarction or hemorrhage may occur. 3 U: weak; û; brisk or increased. 4 Methylenedioxymethamphetamine = "ecstasy." 5 Causes: dehydration, hyponatremia, carebral edema, cardiovascular complications, hyperthermia, rhabdomyolysis. 6 Myelopathy, polyneuropathy, Guillain-Barré syndrome, and rhabdomyolysis may occur in chronic heroin users. 7 D-lysergic acid diethylamide.

latrogenic Encephalopathies

Neurological side effects of diagnostic studies and therapies must be kept in mind in the clinical decision-making process (risk/benefit analysis) and must be considered in the differential diagnosis of encephalopathy. Such side effects are easily mistaken for neurological dysfunction of another etiology. Examples are listed in Table 52 (p. 389).



Signal ____ attenuation (pons)



Central pontine myelinolysis (sagittal/axial T₁-weighted MRI images)



Inhalation of industrial or household chemicals ("sniffing")

- Ethylene oxide (gas sterilization)
- Lead (children)
- Industrial waste
- Organic solvents (hydrocarbons, ketones, esters, alcohols)
- Organic tin compounds (wood care products, silicone rubber, thermal insulators)
- Pesticides
- Mercury
- Thallium (rat poison)

Encephalopathies caused by industrial toxins



latrogenic encephalopathy



Drugs (behavioral changes)

Neuropathy Syndromes

Disturbances of the peripheral nervous system may be subdivided into those affecting neuronal cell bodies (neuronopathy) and those affecting peripheral nerve processes (peripheral neuropathy). Neuronopathies include anterior horn cell syndromes (motor neuron lesions; p.50) and sensory neuron syndromes (sensory neuronopathy, ganglionopathy; pp.2, 107, 390). Motor neuron diseases are described on p. 304. Peripheral neuropathy is characterized by damage to myelin sheaths (myelinopathy) and/or axons (axonopathy). Neuropathies may affect a single nerve (mononeuropathy), multiple isolated nerves (mononeuropathy multiplex), all peripheral nerves generally (polyneuropathy), or all peripheral nerves generally with accentuation of one or a few (focal polyneuropathy). Polyneuropathy may be accompanied by autonomic dysfunction (p. 140). The terms polyneuropathy (PNP) and peripheral neuropathy are often used synonymously. Radiculopathies (nerve root lesions) are classified as either monoradiculopathies or polyradiculopathies, depending on whether a single or multiple roots are involved.

Symptoms and Signs

Peripheral neuropathy causes sensory, motor, and/or autonomic dysfunction. Its etiological diagnosis is based on the pattern and timing of clinical manifestations (Table 53, p. 390).

Sensory dysfunction (p. 106) is often the first sign of neuropathy. Sensory deficits have distinctive patterns of distribution: they may be predominantly proximal or distal, symmetrical (stocking/glove distribution) or asymmetrical (multiple mononeuropathy), or restricted to individual nerves (cranial nerves, single nerves of the trunk or limbs; p. 32 f). Disordered sensory processing (p. 108 f) can produce hyperalgesia (more pain than normal upon noxious stimulation), hyperesthesia (increased tactile sensation with lowering of threshold), paresthesia (spontaneous or provoked abnormal sensation), dysesthesia (spontaneous or provoked, abnormal, painful sensation), or allodynia (pain resulting from nonnoxious stimuli). Damage to rapidly conducting, thickly myelinated A-β fibers causes paresthesiae such as tingling, prickling ("pins and needles"), formication, and sensations of tension, pressure, and swelling. Damage to slowly conducting, thinly myelinated A-δ and C fibers (*small fiber neuropathy*) causes hypalgesia or analgesia with thermal hypesthesia or anesthesia, abnormal thermal sensations (cold, heat), and pain (burning, cutting, or dull, pulling pain).

Motor dysfunction (p. 50). Weakness usually appears first in distal muscles. In very slowly progressive neuropathies, muscles may become atrophic before they become weak, but weakness is usually the initial symptom, accompanied by hyporeflexia or areflexia. The cranial nerves can be affected. Hyperactivity in motor $A-\alpha$ fibers produces muscle spasms, fasciculations, and/or myokymia.

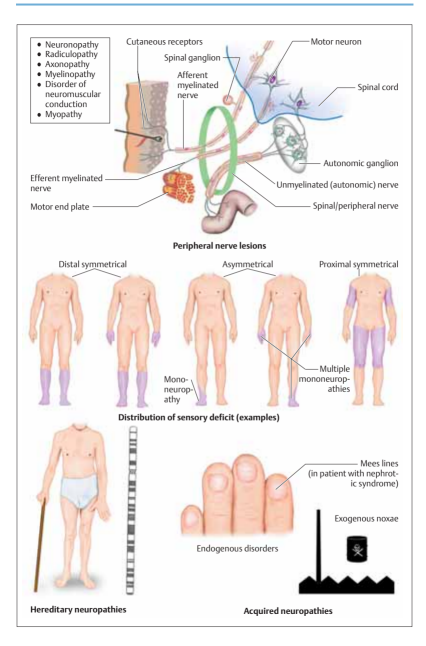
Autonomic dysfunction (p. 146 f) can be manifest as vasomotor disturbances (syncope), cardiac arrhythmias (tachycardia, bradycardia, fixed heart rate), urinary and gastrointestinal dysfunction (urinary retention, diarrhea, constipation, gastroparesis), sexual dysfunction (impotence, retrograde ejaculation), hyperhidrosis or hypohidrosis, pupillary dysfunction, and trophic lesions (skin ulcers, bone and joint changes).

Etiology (Table 54, p. 390)

Polyneuropathies can be hereditary or acquired (see Table 54).

Diagnosis (Table 55, p. 391)

The diagnosis of a neuropathy is based on the characteristic clinical findings and patient history. Additional diagnostic studies not indicated on the basis of the patient history and clinical findings may produce not only unjustified costs but also confounding data, leading occasionally to misdiagnosis. Studies to be performed as indicated include neurophysiological tests (nerve conduction studies, electromyography), laboratory tests (blood, CSF), tissue biopsy (nerve, skin, muscle), and genetic tests.



Radicular Lesions

Symptoms and Signs

Patients usually complain mainly of positive sensory symptoms (tingling, burning, intense pain), which, like the accompanying sensory deficit (mainly hypalgesia, see p. 104 f), are in a dermatomal distribution (p. 32 ff). Weakness, if any, is found mainly in muscles that are largely or entirely innervated by a single nerve root (pp. 32, 50); loss of the segmental deep tendon reflex (p. 40) is, however, a typical early finding. Monoradiculopathy does not cause any evident autonomic dysfunction in the limbs. Lumbar monoradiculopathy is frequently caused by lumbar disk herniation with secondary root compression; typical findings in such cases include exacerbation of radicular pain by coughing, straining at stool, sneezing, or vibration (\Rightarrow the patient adopts an antalgic posture), as well as Lasègue's sign (radicular pain on passive raising of the leg with extended knee) and Bragard's sign (radicular pain on dorsiflexion of the foot with the leg raised and extended). Bladder, bowel, and sexual dysfunction may be caused by a lesion affecting multiple roots of the cauda equina (p. 48), or by processes affecting the spinal cord (pp. 48, 282) or sacral plexus (see below).

Pseudoradicular syndromes (including so-called myofascial syndrome, tendomyalgia, myotendinosis) are characterized by limb pain, localized muscle tenderness, and muscle guarding and disuse, without radicular findings.

Causes

See Table 56, p. 392, and p. 320.

Plexopathy (p. 321)

For clinical purposes the brachial plexus (p. 34) located behind the clavicle may be divided into *supraclavicular* and *infraclavicular* regions. The supraclavicular plexus consists of the primary (ventral and dorsal) roots, mixed spinal nerves, five anterior primary rami, and three trunks; the infraclavicular plexus is composed of the three cords and the terminal nerves. Lesions affecting the supraclavicular plexus can either be preganglionic (intradural, inside the spinal canal) or infraganglionic (extradural, extraforaminal, out-

side the spinal canal). Supraclavicular plexopathies are more common than infraclavicular ones.

Symptoms and Signs

Brachial plexus. Lesions affecting the entire brachial plexus cause anesthesia and flaccid paralysis of the entire upper limb, with muscle atrophy. Lesions of the upper brachial plexus (C5-C6) cause weakness of shoulder abduction and external rotation, elbow flexion, and supination, with preservation of hand movement (Erb palsy). The limb hangs straight down with the hand pronated. A sensory deficit may be found on the lateral aspect of the arm and forearm. Lesions of the lower brachial plexus (C8-T1) mainly cause weakness of the hand muscles (Klumpke-Dejerine palsy); atrophy of the intrinsic muscles produces a claw hand deformity. A sensory deficit is found on the ulnar aspect of the forearm and in the hand. Concomitant involvement of the cervical sympathetic pathway produces Horner syndrome. Erb palsy is more likely to recover spontaneously than Klumpke-Dejerine palsy.

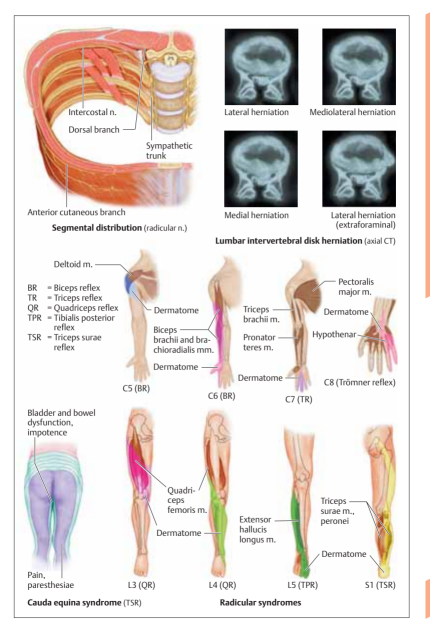
Lumbosacral plexus. Lesions of the *lumbar* plexus (L1–L4) cause weakness of hip flexion and knee extension (as in a femoral nerve lesion) as well as thigh adduction and external rotation. A sensory deficit is found in the affected dermatomes (p. 36). Lesions of the sacral plexus (L5–S3) cause weakness of the gluteal muscles, hamstrings, and plantar and dorsiflexors of the foot and toes. A sensory deficit is found on the dorsal aspect of the thigh, calf, and foot. Lesions of the *lumbar sympathetic trunk* cause leg pain and an abnormally warm foot with diminished sweating on the sole.

Causes

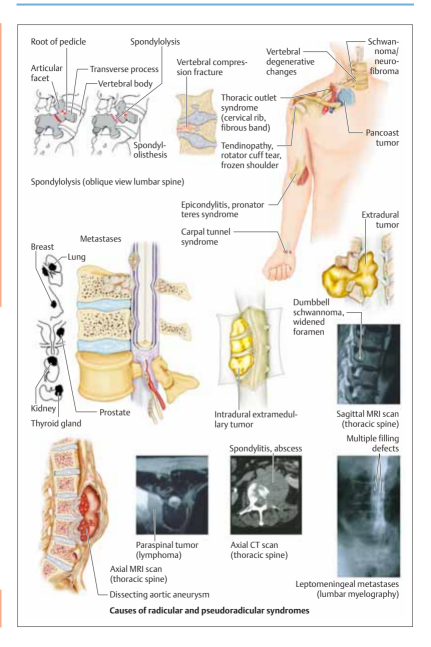
See Table 57, p. 393.

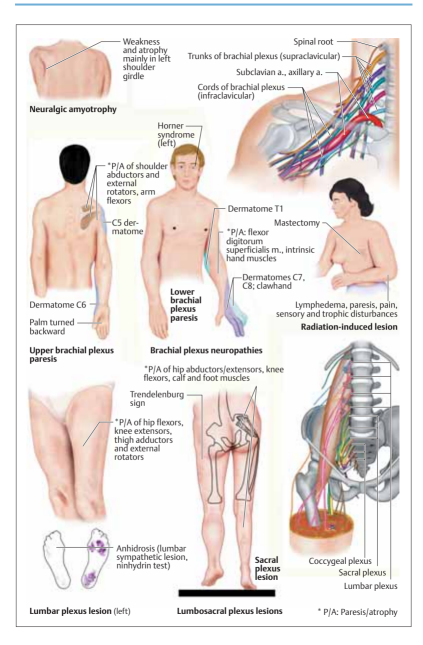
Mononeuropathies (p. 322 f)

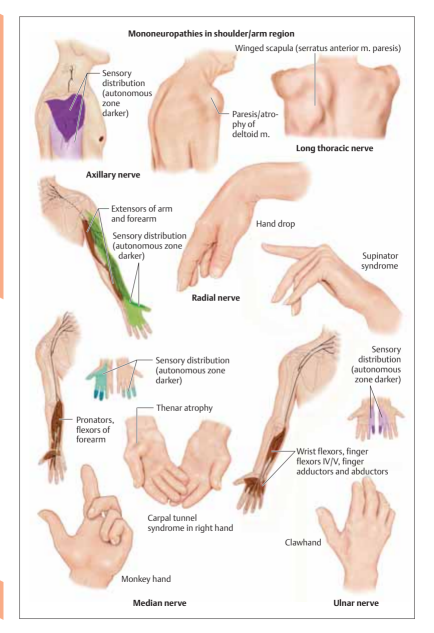
Lesions affecting a single nerve tend to occur at certain favored sites and are usually of mechanical origin (compression, hyperextension, transection) (Table 58, p. 394).

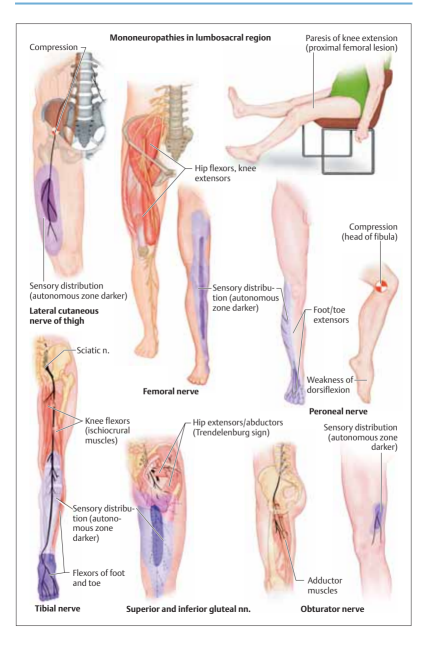


Peripheral Neuropathies









Diabetic Neuropathies

Diabetes Mellitus

Diabetes mellitus (DM) is a syndrome of impaired carbohydrate metabolism due to insulin deficiency. In Type 1 DM (about 10% of cases), the insulin-secreting pancreatic cells are destroyed by an autoimmune process; Type 2 DM (the remaining 90%) is a nonautoimmune disorder typified by insulin resistance and abnormally low insulin secretion, usually in conjunction with obesity. Sequelae of DM include arteriosclerosis, microangiopathy, retinopathy, nephropathy, and peripheral neuropathy. The fasting blood glucose concentration is elevated $(\geq 126 \text{ mg/dl})$, or else the blood glucose concentration is elevated after a standardized oral glucose load. An integrated index of elevated blood glucose concentration over time can be obtained by measuring the concentration of glycosylated hemoglobins (HBA₁, HBA₁ $rac{} \Rightarrow$ 4–6 weeks) and proteins (fructosamine \Rightarrow 8–14 days).

Syndromes

Pathogenesis. Distal symmetric polyneuropathy, a form of diabetic polyneuropathy (DPN), is due to generalized peripheral nerve damage. It is a complication of continuous hyperglycemia and the related metabolic changes (⇒ polyols, phospholipids, fatty acids, oxidative radicals, lack of nerve growth factors). Normalization of the blood glucose concentration by medical treatment can prevent DPN (at least partially) but, long-standing severe DPN, once established, cannot be completely reversed by euglycemia. Pathological examination reveals extensive axon loss, which is thought to be due either to the chronic hyperglycemia itself or to the resulting (perhaps inflammatory) microvascular changes. Repeated episodes of hypoglycemia (p. 308) can also cause neuropathy.

Symptoms and signs. DPN produces both negative neurological signs (sensory loss, sensory ataxia, thermanesthesia, hypalgesia, autonomic dysfunction, paresis) and positive neurological signs (paresthesia, dysesthesia, pain). The manifestations of DPN are classified in Table 59 (p. 395). They may be present in varying combinations.

Diagnosis. DPN is diagnosed in diabetic patients with distal, symmetric, sensorimotor poly-

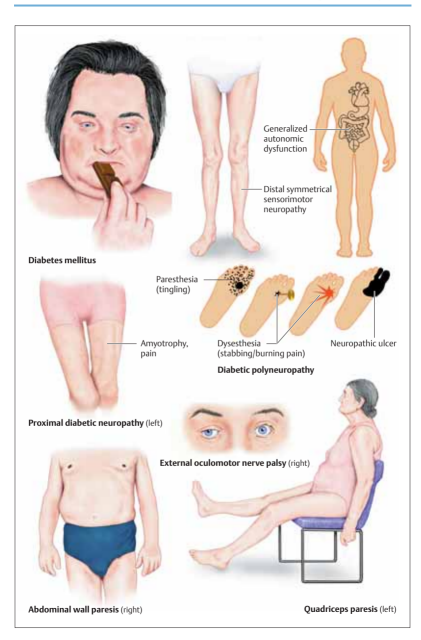
neuropathy of the lower limbs, after the exclusion of other causes (e. g., diabetic lumbosacral or radicular lesions (\Rightarrow Table 59), other neuropathies or primary illnesses); it is usually accompanied by diabetic retinopathy or nephropathy of comparable severity. Other neuropathic syndromes found in diabetes (some symmetric, some asymmetric) require the use of specialized tests for their differential diagnosis (p. 391).

Treatment. The main objective of treatment is normoglycemia. Pain (p. 108) due to diabetic neuropathy usually responds to tricyclic antidepressants (amitryptiline, clomipramine), anticonvulsants (carbamazepine, gabapentin, lamotrigine), antiarrhythmics (lidocaine, mexiletine), capsaicin (administered locally as a 0.075% cream), or transdermal clonidine. Autonomic dysfunction of various types is treated symptomatically. Other factors that may worsen the neuropathy should be avoided (alcohol, vitamin deficiency, medication side effects). The complications of DPN (diabetic foot ulcer, infection, weakness, falls) may require specific treatment.

Uremic Neuropathy

Uremic neuropathy is a distal, symmetrical, sensorimotor, axonal peripheral neuropathy that mainly affects the legs. Paresthesiae and a "restless legs" sensation are typical. Uremic neuropathy may complicate renal failure of any etiology and is treated by therapy of the underlying disease.

Peripheral Neuropathies



Inflammatory Polyneuropathies

Guillain-Barré Syndrome

The term Guillain–Barré syndrome (GBS) covers a group of monophasic, acute, inflammatory polyneuropathies of autoimmune pathogenesis (Table 60, p. 395). Their onset is 1–4 weeks after a respiratory or gastrointestinal infection in two-thirds of all cases. The causative organism often cannot be identified. GBS is known to be associated with certain viruses (cytomegalovirus, Epstein–Barr virus, varicella-zoster virus, HIV \Rightarrow lymphocytic pleocytosis in CSF), bacteria (*Campylobacter jejuni, Mycoplasma pneumoniae*), and vaccines (rabies).

Pathogenesis. The organisms causing the preceding infection are thought to induce T-cell autoreactivity; after a latency period of days to weeks, antigen-specific T and B cells are activated. The target antigen is still unknown. IgG antibodies of various types, produced by the B cells, can be detected in serum in varying concentrations. These antibodies may block impulse conduction (acute paralysis) or activate complement and macrophages (⇒ myelin lesions). T_H1 lymphocytes release proinflammatory cytokines (IFN- γ , TNF- α ; p. 220) that stimulate macrophages (⇒ peripheral nerve lesions). Once the inflammatory response has regenerative subsided, processes (axonal growth and remyelination) begin.

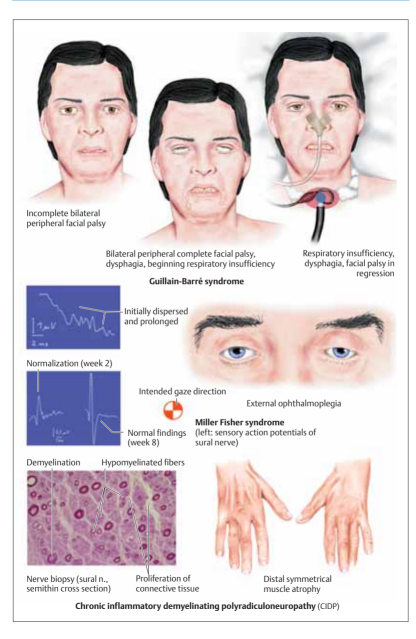
Symptoms and signs. GBS classically presents with an acute ascending and often rapidly progressive symmetrical weakness, areflexia, and relatively mild sensory abnormalities (paresthesiae). Pain is not uncommon, especially at onset; it is often in the back, of shocklike, tingling, aching, or myalgic quality, and may be misattributed to a herniated disk, "the flu," or "rheumatism." Cranial nerve deficits (VII, often bilateral; III, IV, VI, IX, X) are almost always present. So, too, are respiratory weakness and autonomic disturbances (bradycardia or tachycardia, hypotension or hypertension, abnormalities of fluid and electrolyte balance), all of which frequently cause complications. The sudden onset of disease with severe, ascending weakness is often a terrifying experience for patients and their families.

The clinical features and course of GBS are highly variable. Predictors of an unfavorable

outcome include age over 60 years, progression to quadriplegia within one week, the need for mechanical ventilation, and a reduction of the amplitude of motor evoked potentials to less than 20% of normal. The manifestations of less common forms of GBS are listed on p. 395.

Diagnosis (Table 61, p. 396). GBS is diagnosed from its typical clinical features. Neurophysiological findings are used to support the diagnosis, rule out alternative diagnoses, and document the type and extent of peripheral nerve damage. CSF studies are mainly useful for the exclusion of alternative diagnoses. It may be difficult to determine which specific type of GBS is present.

Treatment. Complications of GBS are mainly due to autonomic dysfunction, respiratory insufficiency, and immobility (⇒ deep venous thrombosis, pulmonary embolism, compression neuropathies, pressure sores, contractures). Patients should thus be closely monitored in an intensive care unit, especially in the acute phase. They and their relatives should be offered clear and ample information about the disease, as well as psychological counseling. GBS may be treated by intravenous gammaglobulin therapy or plasmapheresis.



Chronic Inflammatory Demyelinating Polyradiculoneuropathy (CIDP)

CIDP differs from GBS in that it is of subacute onset (slow progression over 2 months or more) and responds readily to immune suppression (corticosteroids, azathioprine, cyclophosphamide; dose titrated to response) in combination with immunoglobulins or plasmapheresis. Its course is usually progressive or relapsing rather than monophasic. It is less commonly preceded by systemic infection than GBS but otherwise has very similar clinical features. Pain may accompany or precede an exacerbation of the illness. Neurophysiological studies reveal evidence of demyelination. The CSF protein concentration is markedly elevated, and sural nerve biopsy reveals chronic demyelination and remyelination with rare inflammatory infiltrates.

Multifocal Motor Neuropathy (MMN)

MMN is characterized by progressive asymmetrical weakness, which usually begins in the upper limbs. The underlying lesion is usually in isolated peripheral nerves (most often radial, median, ulnar, and common peroneal). Muscle spasms and fasciculations are common. Sensory loss, if any, is mild, and muscle atrophy is mild or absent even if weakness is marked. The reflexes may be absent, diminished, or (rarely) brisk. Nerve conduction studies reveal a motor conduction block. There may be an elevated serum concentration of IgM antibodies to GM1. Repeated intravenous administration of immunoglobulin or cyclophosphamide is an effective treatment. The differential diagnosis includes amyotrophic lateral sclerosis, distal spinal muscular atrophy (p. 385), and CIDP.

Paraproteinemic Polyneuropathies

These disorders are most commonly due to nonmalignant monoclonal gammopathies, usually of IgM type, rarely IgG or IgA, though there is progression to plasmocytoma or Waldenström macroglobulinemia in some 20% of cases. The main features of monoclonal gammopathy of undetermined significance (MGUS) include: κ-type, M protein <25 g/l, Bence Jones proteinuria (rare), no skeletal or organ involvement, normal blood smear. The clinical manifestations are of slowly progressive, distal, symmetrical, sensorimotor polyneuropathy, which is sometimes painful. Serum antibodies to myelin-associated glycoprotein (MAG) may be present, and the CSF protein concentration may be elevated. The treatment is by immune suppression, but the ideal type of agent, timing, and dosage have not yet been determined. *POEMS syndrome* (PNP + organomegaly + endocrinopathy + M protein + skin changes; *Crow-Fukase syndrome*) is a rare systemic manifestation of osteosclerotic myeloma.

Neuropathy of Infectious Origin

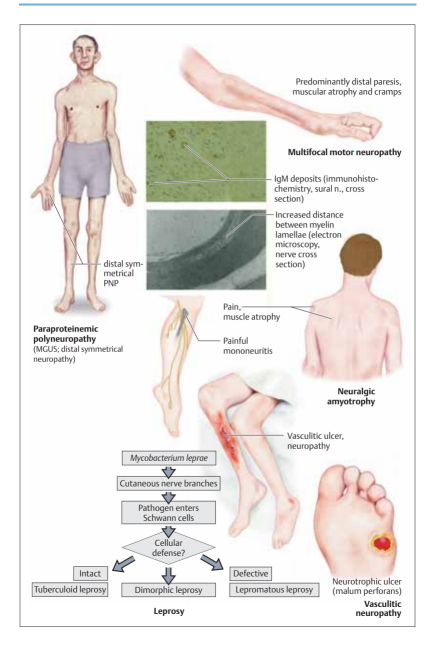
Leprosy, HIV infection, herpes zoster infection, borreliosis, tetanus, botulism, diphtheria, or other infectious diseases may cause neuropathy. Leprosy is the most common cause of peripheral neuropathy around the world. Its pathogenic organism (*Mycobacterium leprae*) attacks peripheral nerves in the cooler parts of the body, such as the skin, nose, anterior portion of the eye, and testes. There is segmental thickening of peripheral nerves (elbow, wrist, ankle). Areas of skin become depigmented and anhidrotic, with dissociated sensory loss. There are different types of leprosy, each of which is associated with a characteristic type of neuropathy.

Neuralgic Amyotrophy

This disorder involves acute, usually nocturnal attacks of severe pain in the shoulder for several days or weeks, followed by weakness and muscle atrophy. Sensory deficits are rare (axillary nerve distribution). The symptoms usually resolve spontaneously.

Vasculitic Neuropathy

Peripheral neuropathy due to connective tissue disease is usually multifocal, rarely symmetric (p. 180). Early treatment by immune suppression improves the outcome. Various connective tissue diseases can produce an isolated sensory trigeminal neuropathy.



Peripheral Nerve Injuries

Peripheral nerves can be temporarily or permanently damaged by pressure, transection, crushes, blows, or traction.

Type of Lesion	Selected Causes/Features	Classification ¹ /Prognosis
Local conduction block, with nor- mal conduction distal to the le- sion	Local demyelination due to com- pression Conduction blockade without EMG evidence of degeneration	Neurapraxia Resolution within a few weeks in most cases
Damage to axon and myelin sheath with preservation of en- veloping structures (Schwann cell basal membrane and en- doneurium); wallerian degenera- tion distal to the lesion	Crushing of nerve Regeneration occurs from proxi- mal to distal along the enveloping structures, taking weeks, months, or years, depending on whether the damage is partial or com- plete ²	Axonotmesis EMG evidence of reinnervation is seen first in muscle groups proxi- mal to the lesion, and later in dis- tal groups
Damage to axon, myelin sheath, and enveloping structures; wal- lerian degeneration distal to the lesion	Excessive traction, open incision wound Axon regeneration greatly limited; anomalous regeneration and neu- roma development are common	Neurotmesis Full recovery is unusual

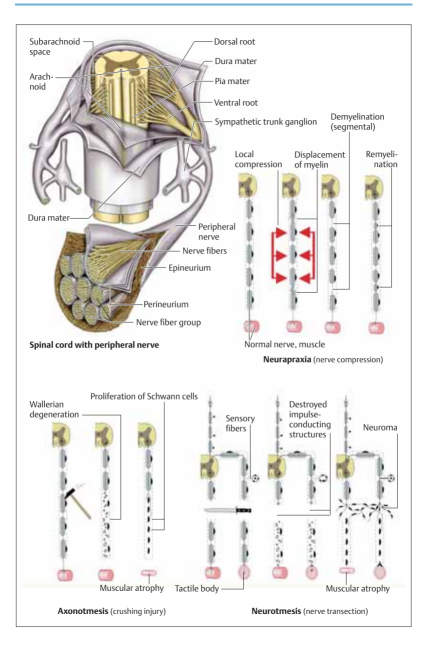
1 Seddon (1943). 2 Axons regenerate at 1–2 mm/day proximally, more slowly distally.

Pathogenesis

Local nerve compression displaces the axoplasm laterally from the site of compression. This causes invagination and subsequent demyelination at the nodes of Ranvier, so that saltatory impulse conduction is blocked. Compression preferentially impairs conduction in large-caliber fibers. *Crushing of a nerve* destroys the axoplasm but not the basal lamina. Schwann cells and axon processes regenerate in the damaged region and distally along the intact enveloping structures until they reach the effector muscle. *Nerve transection* is followed by axonal and Schwann cell proliferation, which may lead to the formation of a neuroma at the proximal nerve stump. Suturing the proximal and distal stumps together enables the regenerating fibers to enter the distal enveloping structures and regenerate further, but the function of the nerve is usually not fully restored to its original state.

Treatment

Type of Lesion	Treatment
Nerve root avulsion Brachial plexus injury	 Surgery (e. g., tenodesis, tendon-muscle transfer), treatment of pain Open ⇒ primary nerve suture Closed ⇒ surgical exploration if there is no reinnervation in 3–5 months; if function fails to improve, other surgical procedures for restoration of function can be considered
Neurapraxia or axonotmesis (nontransecting injury)	 Diagnosed from clinical findings, EMG, and nerve conduction studies at presentation and 3 weeks later; treated with physical therapy Clinical and neurophysiological re-assessment every 2–5 months Clinical and neurophysiological improvement ⇒ further physical therapy No clinical or neurophysiological improvement ⇒ corrective surgery 2–3 weeks after local injury (e. g., gunshot wound) or 4–5 months after extensive injury (e. g., traction injury)
Neurotmesis (nerve transection)	 Primary suture of nerve cut by knife, glass, etc. Secondary suture 2–4 weeks after crushing injury and/or destruction of epineurium



Nonmetabolic Hereditary Neuropathies (Tables 62 and 63, p. 396 f)

For neuropathies associated with systemic disease, see p. 280.

Hereditary Motor–Sensory Neuropathy (HMSN)

These are the most common among the hereditary neuropathies, all of which are rare.

HMSN type I is characterized by high pedal arches (pes cavus), hammer toes (digitus malleus), distal weakness and atrophy, loss of vibration sense with preservation of position sense, areflexia, and unsteady gait (p. 60; frequent stumbling, steppage gait). Some peripheral nerves are palpably thickened in half of the cases (e.g., the greater auricular, ulnar, or common peroneal nerve), and tremor is present in one-third. The clinical picture is highly variable. The type I phenotype is produced by three different genotypes: CMT1 (autosomal dominant), CMT4 (autosomal recessive), and CMTX (X-linked).

HMSN type II. CMT2A and B resemble HMSN type I but begin later, with only rare areflexia and no palpable thickening of peripheral nerves. CMT2C is characterized by vocal cord paralysis (hoarseness, inspiratory stridor), proximal and distal weakness, distal muscle atrophy, and areflexia.

HMSN type III. This rare polyneuropathy (eponym: Dejerine–Sottas disease) becomes manifest at birth or in childhood with generalized weakness, areflexia, and palpable nerve thickening. Hearing loss, skeletal deformity, and sensory deficits (⇔ataxia) ensue as the disease progresses.

Hereditary Neuropathy with Pressure Palsies (HNPP)

HNPP is characterized by recurrent, transient episodes of weakness and sensory loss after relatively mild compression of a peripheral nerve (ulnar, peroneal, radial, or median nerve). There may be evidence of a generalized polyneuropathy or a painless plexopathy. The "sausagelike" pathological changes seen on sural nerve biopsy are the origin of the alternative name, tomaculous neuropathy (from Latin tomaculum, "sausage").

Metabolic Hereditary Neuropathies

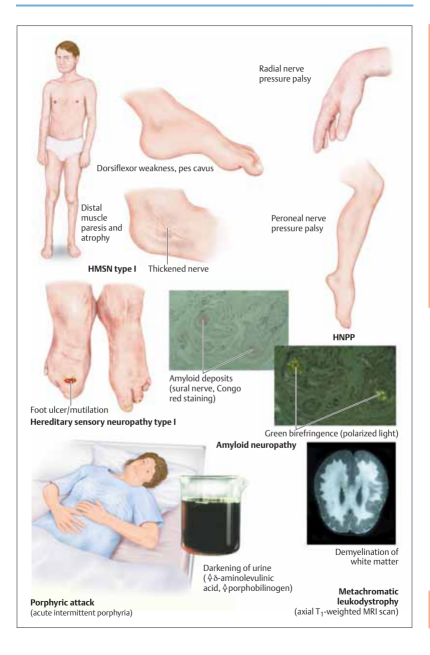
Other members of this class are listed in the section on metabolic diseases (p. 306 ff).

Porphyria

Among the known porphyrias, four hepatic types are associated with encephalopathy and peripheral neuropathy: variegate porphyria, acute intermittent porphyria, hereditary coproporphyria, and δ -aminolevulinic acid dehydrase deficiency (autosomal recessive; the others are autosomal dominant). The porphyrias are hereditary enzymopathies affecting the biosynthesis of heme. Severe peripheral neuropathy is seen during attacks of acute porphyria, which are most often precipitated by medications and hormonal influences (also fasting, alcohol, and infection). The manifestations of porphyria include colicky abdominal pain, pain in the limbs, paresthesiae, tachycardia, and variable degrees of weakness. Encephalopathy is manifest as confusion, lack of concentration, somnolence, psychosis, hallucinations and/or epileptic seizures. The diagnosis of porphyria is based on the demonstration of porphyrin metabolites in the urine and feces.

Neuropathy Due to Hereditary Disorders of Lipid Metabolism

Polyneuropathy occurs in metachromatic leukodystrophy Krabbe (p. 306), disease (p. 307), abetalipoproteinemia (p. 300), adrenomyeloneuropathy (p. 384), Tangier disease (tonsillar hypertrophy, hepatosplenomegaly, low serum cholesterol, serum HDL deficiency), Fabry disease (punctate red angiokeratoma on buttocks and in the genital and periumbilical areas, retinovascular changes, corneal deposits, nephropathy, painful neuropathy; glycosphingolipid deposition due to a-galactosidase deficiency), and Refsum disease. The last is an autosomal recessive disorder of phytanic acid metabolism in which phytanic acid accumulation leads to tapetoretinal degeneration, night blindness, and a distal, symmetric polyneuropathy with peripheral nerve thickening. The CSF protein concentration is markedly elevated, but the CSF cell count is normal. The serum phytanic acid concentration is elevated.



Myopathic Syndromes

Myopathies are diseases of muscle. Many different hereditary and acquired diseases attack muscle, sometimes in combination with other organs. The diagnosis and classification of the myopathies have been transformed in recent years by the introduction of molecular biological tests for the hereditary myopathies, but their treatment remains problematic. The management of the hereditary myopathies currently consists mainly of genetic counseling and the attempt to provide an accurate prognosis.

Symptoms and Signs (Table 64, p. 397)

Weakness (p. 52) is the most common sign of myopathy; it may be of acute, rapidly progressive, or gradual onset, fluctuating, or exerciseinduced. It may be local (restricted to the muscles of the eye, face, tongue, larynx, pharynx, neck, arms, legs, or trunk), proximal, or distal, asymmetric or symmetric. Myalgia, muscle stiffness, and muscle spasms are less common. There may be muscle atrophy or hypertrophy, often in a typical distribution, whose severity depends on the type of myopathy. Skeletal deformity and/or abnormal posture may be a primary component of the disease or a consequence of weakness. Other features include acute paralysis, myoglobulinemia, cardiac arrhythmia, and visual disturbances.

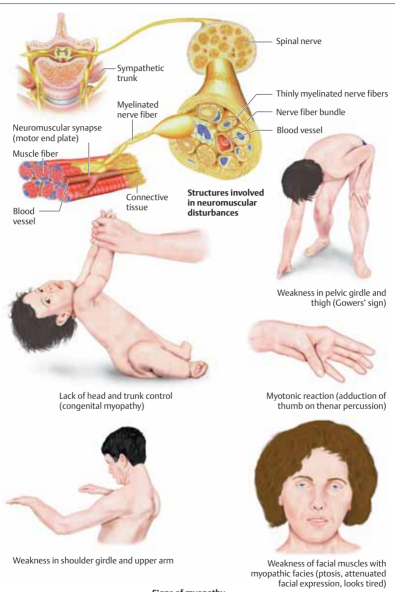
Causes

For a list of causes of hereditary and acquired myopathies, see Tables 65 and 66, p. 398.

Diagnosis (Table 67, p. 399)

The myopathies are diagnosed primarily by history and physical examination (p. 52). Pharmacological tests are used for the differential diagnosis of myasthenia. Neurophysiological studies are used to rule out neuropathy (p. 391), to determine the specific type of acute muscle change, or to identify disturbances of muscular impulse generation and conduction. Various laboratory tests are helpful in myopathies due to biochemical abnormalities; imaging studies of muscle aid in the differential diagnosis of atrophy and hypertrophy. Muscle biopsy is often needed for a definitive diagnosis. Molecular biological studies are used in the diagnosis of hereditary myopathies.

Myopathies



Signs of myopathy

Muscular Dystrophies

The muscular dystrophies—myopathies characterized by progressive degeneration of muscle are mostly hereditary.

■ Pathogenesis (Table 65, p. 398)

Dystrophinopathies are X-linked recessive disorders due to mutations of the gene encoding dystrophin, a protein found in the cell membrane (sarcolemma) of muscle fibers. Such mutations cause a deficiency, alteration, or absence of dystrophin. The functional features of dystrophin are not fully understood; it is thought to have a membrane-stabilizing effect. Some forms of limb girdle dystrophy (e.g., sarcoglycanopathy) are due to mutation of genes encoding dystrophin-associated glycoproteins, while others are due to mutation of genes encoding intracellular enzymes such as calpain-3. Emery-Dreifuss muscular dystrophy is due to a mutation of the gene for emerin, a nuclear membrane protein whose exact function is unknown.

Symptoms and Signs

Muscular dystrophies may be characterized by atrophy, hypertrophy, or pseudohypertrophy and are further classified by their mode of inheritance, age of onset, and distribution. Other features such as myocardial involvement, contractures, skeletal deformity, endocrine dysfunction, and ocular manifestations may point to one or another specific type of muscular dystrophy. Each type has a characteristic course (Table 68, p. 400).

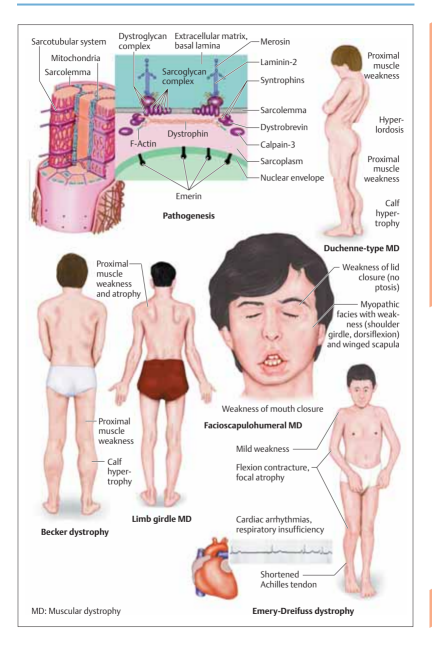
Diagnosis

The history and physical examination are supplemented by additional diagnostic studies including ECG, creatine kinase fractionation (CK-MM), EMG, and DNA studies. If DNA analysis fails to reveal a mutation, immunohistochemical techniques, immune blotting, or the polymerase chain reaction can be used to detect abnormalities of dystrophin and sarcoglycan (e.g., in muscle biopsy samples) and thereby distinguish between Duchenne and Becker muscular dystrophy, or between dystrophinopathies and other forms of muscular dystrophy. DNA tests are used for the identification of asymptomatic female carriers (in whom muscle biopsy is hardly ever necessary), and for prenatal diagnosis.

Treatment

The goal of treatment is to prevent contracture and skeletal deformity and to keep the patient able to sit and walk for as long as possible. The patient's diet should be monitored to prevent obesity. The most important general measures are genetic counseling, social services, psychiatric counseling, and educating the patient on the special risks associated with general anesthesia. The type of schooling and employment must be appropriately suited to the patient's individual abilities and prognosis. Physical therapy includes measures to prevent contractures, as well as breathing exercises (deep breathing, positional drainage, measures to counteract increased inspiratory resistance). Patients with alveolar hypoventilation may need intermittent ventilation with continuous positive airway pressure (CPAP) at night. Orthoses may be helpful, depending on the extent of weakness (night splints to prevent talipes equinus, seat cushions, peroneal springs, orthopedic corsets, leg orthoses). Home aids may be needed as weakness progresses (padding, eating aids, toilet/bathing aids, stair-lift, mechanized wheelchair, specially adapted automobile). Surgery may be needed to correct scoliosis, prevent contracture about the hip joint (iliotibial tract release), and correct winging of the scapula (scapulopexy/scapulodesis) and other deformities and contractures. Intracardiac conduction abnormalities (e.g., in Emery-Dreifuss muscular dystrophy) require timely pacemaker implantation. Heart transplantation may be needed when severe cardiomyopathy arises in conjunction with certain types of muscular dystrophy (Becker, Emery-Dreifuss; Table 68, p. 400).

Myopathies



The Myotonias (Table 69, p. 401)

Pathogenesis

Point mutations in ion channel genes cause channel defects that render the muscle cell membrane electrically unstable (Table 65, p. 398), leading to involuntary muscle contraction.

Symptoms and Signs

The transient, involuntary muscle contractions are perceived as stiffness. Depolarizing muscle relaxants used in surgery can trigger severe myotonia in susceptible patients. Acute, generalized myotonia can also be induced by tocolytic agents such as fenoterol.

Diagnosis

Myotonia is diagnosed from the observation of involuntary muscle contraction after voluntary muscle contraction (action myotonia) or percussion (percussion myotonia), along with the characteristic EMG findings. Specific forms of myotonia are diagnosed by their mode of inheritance and clinical features, and molecular genetic analysis. The serum creatine kinase concentration is usually not elevated, and there is usually no muscle atrophy, except in myotonic dystrophy. Muscle hypertrophy is present in myotonia congenita. Myotonic cataract is found in myotonic dystrophy and proximal myotonic myopathy; slit-lamp examination is indicated in patients with these disorders.

Treatment

Membrane-stabilizing drugs such as mexiletine alleviate myotonia; cardiac side effects may be problematic, particularly in myotonic dystrophy. Cold exposure should be avoided.

Episodic Paralyses (Table 69, p. 401)

Pathogenesis

Hyperkalemic and normokalemic paralysis, potassium-aggravated myotonia (PAM = myotonia fluctuans), and paramyotonia congenita are due to *sodium channel* dysfunction, while hypokalemic paralysis is due to *calcium channel* dysfunction.

Symptoms and Signs

In hypokalemic and hyperkalemic myotonia, there are irregularly occurring episodes of flaccid paresis of variable duration and severity, with no symptoms in between. The anal and urethral sphincters are not affected. In *paramyotonia congenita*, muscle stiffness increases on exertion (paradoxical myotonia) and is followed by weakness. Cold exposure worsens the stiffness.

Diagnosis

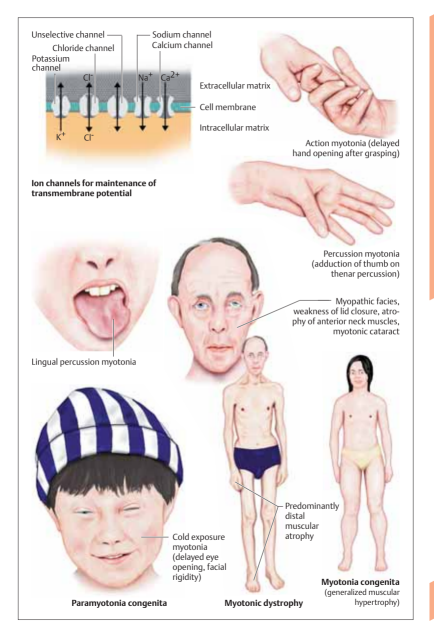
The diagnosis can usually be made from the personal and family history, abnormal serum potassium concentration, and molecular genetic findings (mutation of the gene for a membrane ion channel). If the diagnosis remains in question, provocative tests can be performed between attacks. The induction of paralytic attacks by administration of glucose and insulin indicates hypokalemic paralysis, while their induction by potassium administration and exercise (e.g. on a bicycle ergometer) indicates hyperkalemic paralysis. The diagnosis of paramyotonia congenita is based on the characteristic clinical features (paradoxical myotonia, exacerbation by cold exposure), autosomal dominant inheritance, and demonstration of the causative point mutation of the sodium channel gene.

Treatment

Acute attacks. Milder episodes of weakness in *hypokalemic* disorders need no treatment, while more severe episodes can be treated with oral potassium administration. Milder episodes of weakness in *hyperkalemic* disorders also need no treatment; more severe episodes may require calcium gluconate i. v., or salbutamol by inhaler.

Prophylaxis. *Hypokalemic paralysis:* Low-salt, low-carbohydrate diet, avoidance of strenuous exercise; oral acetazolamide or spironolactone. *Hyperkalemic paralysis:* high-carbohydrate diet; avoidance of strenuous exercise and cold; oral hydrochlorothiazide or acetazolamide.

Myopathies



Congenital Myopathies

The typical pathological findings on muscle biopsy distinguish this group of disorders both from the congenital muscular dystrophies and from muscle changes secondary to peripheral neuropathy. Some congenital myopathies have distinctive clinical features. Proximal flaccid weakness is usually present at birth (floppy baby); skeletal deformities may also be seen (e. g., high palate, hip luxation, pes cavus, chest deformities). Many congenital myopathies progress slowly, causing little or no disability; the CK and EMG may be only mildly abnormal, or not at all. Some types can be diagnosed by genetic analysis (Table 70, p. 402).

Metabolic Myopathies

In most metabolic myopathies (Table 71, p. 402 f), exercise induces myalgia, weakness, and muscle cramps, and myoglobinuria. Progressive proximal weakness is seen in myopathy due to acid-maltase deficiency (glycogen storage disease type II), debrancher deficiency (glycogen storage disease type III), or primary myopathic carnitine deficiency.

Mitochondrial myopathies. Pyruvate and fatty acids are the most important substrates for mitochondrial ATP synthesis, which occurs by oxidative phosphorylation, a function of the respiratory chain enzymes (found on the inner mitochondrial membrane). β -oxidation occurs in the mitochondrial matrix. The respiratory chain enzymes are encoded by both mitochondrial and nuclear DNA (mtDNA, nDNA).

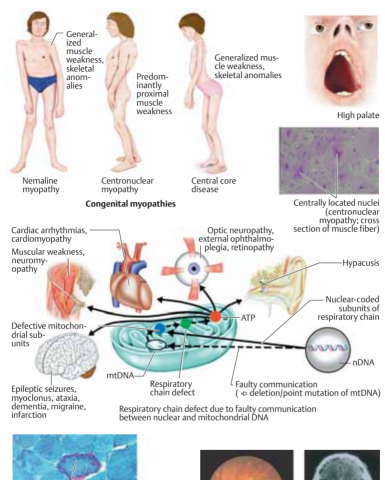
The mitochondrial myopathies are a heterogeneous group of disorders whose common feature is dysfunction of the respiratory chain, β oxidation, or both. These disorders have varying clinical and biochemical features (Table 71, p. 402 f); their inheritance is either maternal or sporadic; non-heritable cases also occur through mtDNA mutations. These disorders may affect multiple organ systems, e.g.:

- Muscle (reduced endurance, pain, cramps, myoglobinuria)
- CNS (seizures, headache, behavioral abnormalities)
- Eye (ptosis, external ophthalmoplegia, tapetoretinal degeneration)

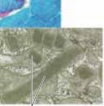
- Ear (hearing loss)
- Heart (arrhythmia, heart failure)
- Gastrointestinal system (diarrhea, vomiting)
- Endocrine system (diabetes mellitus, hypothyroidism)
- ANS (impotence, sweating)

The diagnosis is based on the clinical features, laboratory tests (elevated lactate concentration at rest in serum, sometimes also in CSF, with sustained increase after exercise), muscle biopsy (ragged red fibers, sometimes with cytochrome-c oxidase deficiency), and molecular studies (mtDNA analysis of muscle, platelets, leukocytes). There is no etiological treatment for the mitochondrial myopathies at present; a lowfat, carbohydrate-rich diet is recommended in disorders with defective β-oxidation, and carnitine supplementation in those with systemic carnitine deficiency. Coenzyme Q₁₀, vitamin K₃, vitamin C, and/or thioctic acid supplements are recommended in disorders with impaired respiratory chain function.

Myopathies



Mitochondrial [/] accumulation (ragged red fiber; cross section of muscle fiber)



Paracrystalline mitochondrial inclusions (electron microscopy)



Tapetoretinal degeneration (CPEO)



└── Occipital infarcts (MELAS syndrome, axial CT scan)

Mitochondrial myopathies

Myasthenic Syndromes

Myasthenia Gravis (MG)

Pathogenesis. The exercise-induced weakness that typifies MG is due to impaired transmission at the neuromuscular junction, which is, in turn, due to an underlying molecular lesion affecting the nicotinic acetylcholine receptor (AChR) in the postsynaptic membrane of the muscle cell. Circulating IgG autoantibodies to this receptor impair its function, speed its breakdown, and induce complement-mediated damage to the muscle cell membrane. Recently the anti-MuSK (receptor tyrosine kinase) antibody has been detected in about half of patients who are seronegative for AChR antibodies. The thymus plays an important role in this autoimmune disorder (it is normally a site of maturation and removal of autoreactive T lymphocytes). MG is usually acquired late in life; there are also rare congenital and familial forms.

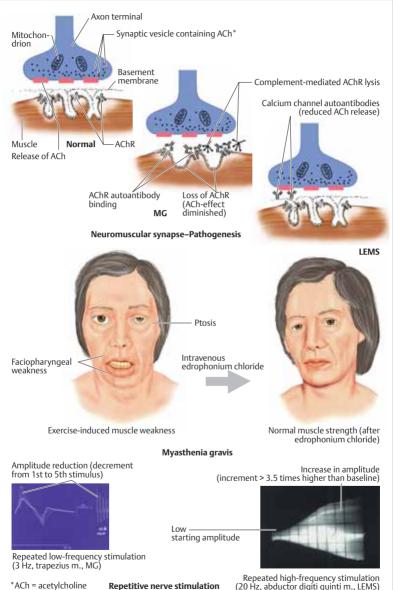
Symptoms and signs. MG is characterized by asymmetric weakness and fatigability of skeletal muscle that worsens on exertion and improves at rest. Weakness often appears first in the extraocular muscles and remains limited to them in some 15% of cases (ocular myasthenia), but progresses to other muscles in the rest (generalized myasthenia). The facial and pharyngeal muscles may be affected, resulting in a blank facial expression, dysarthria, difficulty in chewing and swallowing, poor muscular control of the head, and rhinorrhea. Respiratory weakness leads to impairment of coughing and an increased risk of aspiration. It may become difficult or impossible for the patient stand up, remain standing, or walk, and total disability may ensue. Myasthenia can be aggravated by certain medications (Table 72, p. 403), infections, emotional stress, electrolyte imbalances, hormonal changes, and bright light (eyes), and is often found in association with hyperthyroidism, thyroiditis, rheumatoid arthritis, and connective tissue disease. Myasthenic or cholinergic crises can be life-threatening (Table 73, p. 404). Diagnosis. The diagnosis is based on the characteristic history and clinical findings, supported by further tests that are listed in Table 74 (p. 404).

Treatment. Ocular MG is treated symptomatically with an acetylcholinesterase (AChE) inhibi-

tor, such as pyridostigmine bromide; if the response is insufficient, corticosteroids or azathioprine can be added. Generalized MG is treated initially with AChE inhibitors and, if the response is insufficient, with corticosteroids, azathioprine, intravenous gammaglobulin, or plasmapheresis; once the patient's condition has stabilized, thymectomy is performed. Further treatment depends on the degree of improvement achieved by these measures. The mortality of MG with optimal management is less than 1%. Most patients can lead a normal life but need lifelong immunosuppression. Specific measures are needed to manage respiratory crises, thymoma, and pregnancy in patients with MG, and for the treatment of neonatal, congenital, and hereditary forms of MG.

Lambert–Eaton Myasthenic Syndrome (LEMS)

LEMS is caused by autoantibodies directed mainly against voltage-gated calcium channels in the presynaptic terminal of the neuromuscular junction; diminished release of acetylcholine from the presynaptic terminal is the result. LEMS is often a paraneoplastic manifestation of bronchial carcinoma, sometimes appearing before the tumor becomes clinically evident. It is characterized by proximal (leg) weakness that improves transiently with exercise but worsens shortly afterward. There are also autonomic symptoms (dry mouth) and hyporeflexia. EMG reveals a diminished amplitude of the summated muscle action potential, which increases on high-frequency serial stimulation. The treatment is with 3,4-diaminopyridine (which increases acetylcholine release) and AChE inhibitors. Immune suppression and chemotherapy of the underlying malignancy can also improve LEMS.



Repetitive nerve stimulation

(20 Hz, abductor digiti quinti m., LEMS)

Myositis

The myositides (inflammatory myopathies) are a heterogeneous group of disorders, causing three distinct clinical syndromes: polymyositis (PM), dermatomyositis (DM), and inclusion body myositis (IBM).

Pathogenesis

Most myositides found in the temperate zones are autoimmune diseases of unknown cause, characterized histologically by muscle inflammation and fibrosis and loss of muscle fibers. In PM, cytotoxic CD8⁺ T cells penetrate and damage muscle fibers (⇒ intramuscular cellular infiltrates). CD8+ T cell activation is induced by abnormal expression of class I HLA antigens on the surface of the muscle fibers, which are normally HLA-negative. DM is thought to be largely due to antibodies against blood vessels within muscle, which activate the complement system (membrane attack complex). Vascular endothelial damage ultimately leads to ischemia and death of muscle tissue (\Rightarrow perifascicular atrophy). Inflammatory T cells and macrophages migrate into muscle and cause further damage. IBM is of unknown pathogenesis. Infectious myositis may be due to bacteria, viruses, parasites, or fungi.

Syndromes

Polymyositis (PM) begins with weakness of the proximal muscles of the lower limbs, which then progresses and slowly spreads to the upper limbs. The deltoid and neck flexor muscles are commonly involved. Dysphagia may be present. The involved muscles eventually become atrophic. In overlap syndrome, myositis appears together with another autoimmune disease, e.g., progressive systemic sclerosis, systemic lupus erythematosus, rheumatoid arthritis, polyarteritis nodosa, polymyalgia rheumatica, or Sjögren syndrome. Myalgia is often the major symptoms in patients with PM, as also in patients with hypereosinophilia syndrome (Churg-Strauss syndrome) or eosinophilic fasciitis (Shulman disease).

Dermatomyositis (DM) progresses more rapidly than PM and is distinguished from it mainly by the bluish-red or purple (heliotrope) rash found on exposed areas of the skin (eyelids, cheeks, neck, chest, knuckles, and extensor surfaces of the limbs). Small hemorrhages and telangiectasias are found in the nailbeds; affected children may have subcutaneous calcium deposits. Cancer accompanies DM six times more frequently than PM; DM is also associated with scleroderma and mixed connective tissue disease.

Inclusion body myositis (IBM) is characterized by distal (sometimes asymmetric) weakness and muscle atrophy, mainly in the lower limbs (plantar flexors), with early loss of the quadriceps reflexes. There are both sporadic and hereditary forms of IBM (see also p. 252).

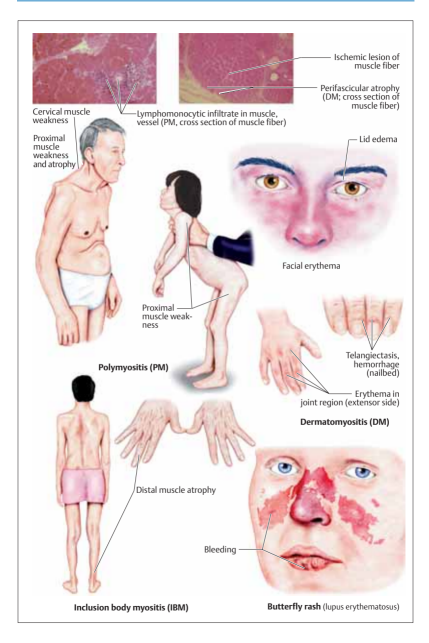
Diagnosis

The myositides are diagnosed by history and physical examination, elevated serum concentration of sarcoplasmic enzymes (particularly CK-MM), and characteristic findings on EMG and muscle biopsy. Muscle atrophy can also be assessed with various imaging techniques (CT, MRI, ultrasonography). The presence of antibodies in association with a connective tissue disease may be relevant to the diagnosis (p. 180).

Treatment

PM and DM are treated by immune suppression, e.g., with corticosteroids, azathioprine, or intravenous gammaglobulin (ivig). Physical therapy is begun once the patient's condition has stabilized. IBM may respond to intravenous immunoglobulin therapy.

Myopathies



Muscle Pain (Myalgia)

Myalgia is an aching, cramping, or piercing pain in muscle. It is triggered by stimulation of nociceptors (p. 108). Pressure or traction on a muscle causes myalgia that subsides once the mechanical stimulus is removed, while inflammatory and other lesions in muscle cause persistent and gradually increasing myalgia. Muscle ischemia and/or metabolic dysfunction are reflected by myalgia occurring only during muscle activity. Myalgia includes *allodynia*, which is defined as pain induced by normally nonpainful stimuli and is explained by the sensitization of nociceptors by pain-related substances such as bradykinin, serotonin, and prostaglandin. A "charleyhorse" is a type of myalgia that normally begins 8–24 hours after muscle overuse (simultaneous stretching and contraction) and lasts 5–7 days. It is caused by an inflammatory reaction to muscle fiber damage. Myalgia can be triggered by disorders whose primary pathology lies anywhere in the nervous system (peripheral nerve, spinal cord, brain).

Causes of Myalgia

Type of Myalgia	Selected Causes
Localized myalgia • Hematoma • Myositis • Ischemic • Toxic-metabolic • Overactivity • Exercise- induced	 Trauma, coagulopathy Infectious: Streptococcal infection, trichinosis, influenza, epidemic pleurodynia. Non-infectious: Nodular focal myositis, eosinophilic fascilitis, sarcoidosis, myositis ossificans Arteriosclerosis (intermittent claudication), embolism Acute alcoholic myopathy, metabolic myopathy (pp. 402, 405) Stiff-man syndrome, neurogenic myotonia, tetanus, strychnine poisoning, amyotrophic lateral sclerosis, tetany Metabolic myopathy, arteriosclerosis, physical exertion
ParkinsonianMuscle spasmPain at rest	 Rigidity Polyneuropathy, metabolic disorder (electrolyte imbalance, uremia, thyroid dysfunction) Restless legs syndrome, painful legs and moving toes syndrome
Generalized myalgia Myositis Toxic-metabolic Other	 Polymyositis/dermatomyositis (p. 344) Hypothyroidism, medications², mitochondrial myopathy (pp. 340, 402, 405) Polymyalgia rheumatica, amyloidosis, osteomalacia, Guillain-Barré syndrome, porphyria, hypothyroidism, corticosteroid withdrawal, fibromyalgia

1 E.g., emetine, lovastatin, and ε-aminocaproic acid.

(Adapted from Layzer, 1994)

Rhabdomyolysis

Local or generalized damage to skeletal muscle can cause myoglobinuria and an elevated serum concentration of creatine kinase, usually accompanied by the acute onset of proximal or diffuse weakness, with myalgia, muscle swelling, and general manifestations including nausea, vomiting, headache, and sometimes fever. The urine may be discolored at the onset of symptoms or several hours later. Rhabdomyolysis can be caused by certain types of myopathy (e. g., polymyositis, central core disease, metabolic myopathies; pp.402, 405), by muscle strain or trauma (long-distance walking or running, heat stroke, delirium tremens, status epilepticus), by toxic substances (see below), and by infectious disease (bacterial sepsis, influenza, coxsackievirus or echovirus infection).

Malignant Hyperthermia (MH)

This life-threatening disorder of skeletal muscle function is characterized by hyperthermia, muscle rigidity, hyperhidrosis, tachycardia, cyanosis, lactic acidosis, hyperkalemia, massive elevation of the serum creatine kinase concentration, and myoglobinuria. It is induced by anesthetic agents such as halothane and succinylcholine. The predisposition to MH is inherited as an autosomal dominant trait (gene loci: 19q13.1, 17q11–24, 7q12.1, 5p, 3q13.1, 1q32). The creatine kinase level may be chronically elevated in susceptible individuals, who can be identified with an in vitro contracture test performed in specialized laboratories. Persons suffering from central core disease, multicore disease, and King–Denborough syndrome (dwarfism, skeletal anomalies, ptosis, high palate) are also at risk for MH. *Treatment:* dantrolene.

Malignant neuroleptic syndrome clinically resembles MH; unlike MH, however, it is usually of subacute onset (days to weeks), it is not hereditary, and it is triggered by psychotropic drugs (haloperidol, phenothiazines, lithium). Malignant neuroleptic syndrome can also be induced by abrupt withdrawal of dopaminergic agents in patients with Parkinson disease.

Toxic Neuromuscular Syndromes

The muscle fiber lesions regress if the responsible substance is eliminated in timely fashion (Table 75, p. 405).

Myopathy in Endocrine Disorders

Hyperthyroidism or hypothyroidism, hyperparathyroidism, Cushing syndrome, steroid myopathy, and acromegaly all cause proximal weakness, while Addison disease and primary hyperaldosteronism usually cause generalized weakness. Timely correction of the endocrine disorder or withdrawal of steroid drugs is usually followed by improvement.

Critical Illness Polyneuropathy (CIP) and Critical Illness Myopathy (CIM)

Sepsis is the most common cause not only of encephalopathy (see p. 312) but also of CIP and CIM. CIP is an acute, reversible, mainly axonal polyneuropathy. It causes distal, symmetric weakness with prominent involvement of the muscles of respiration, resulting in prolonged ventilator dependence and delayed mobilization. CIM causes generalized weakness. The clinical differentiation of CIM and CIP is difficult and often requires muscle biopsy.

Paraneoplastic Syndromes

(Table 76, p. 406)

Distant neoplasms can affect not only the CNS (see p. 388) but also the PNS and skeletal muscle. Remarkably, paraneoplastic syndromes sometimes appear months or years before the underlying malignancy becomes clinically manifest. Paraneoplastic neuromuscular syndromes typically present with marked weakness of subacute onset (i.e., developing over several days or weeks).

A Diagnostic Evaluation

- History and Physical Examination
- Additional Studies

A detailed description of diagnostic evaluation procedures can be found in the textbooks listed on p. 409. The goals of history-taking, physical examination, and additional testing (if necessary) are:

- Data collection (manifestations of disease)
- Localization of the lesion
- Provision of an etiological diagnosis

Data Collection

The diagnostic process begins with the history and physical examination. The history provides information about the patient's experience of illness, the temporal course of symptom development, and potentially relevant familial, social, occupational, and hereditary factors. An inaccurate or incomplete history is a frequent cause of misdiagnosis.

History. The physician engages the patient in a structured conversation about the manifestations of the illness. The physician must remember that the patient is the "expert" in this situation, as the patient alone knows what is troubling him (though perhaps helpful information can also be obtained from a close relative or friend). The physician aims to obtain accurate information on the nature, location, duration, and intensity of the symptoms by listening patiently and asking directed questions in an atmosphere of openness and trust. Questionnaires, computer programs, and ancillary personnel cannot be used for primary history-taking, as they do not enable the construction of a trusting physician-patient relationship (though they may provide useful additional information at a later stage). Some important elements of the case history are as follows.

- Nature of symptoms. The physician must ascertain, by detailed questioning if necessary, that he understands the patient's complaints in the same sense that the patient means to convey. "Blurred vision" may mean diplopia, "dizziness" may mean gait ataxia, "headache" may mean hemicrania, "numbness" may mean paresthesia—but patients may use all of these terms with other meanings as well.
- Severity of symptoms. Quality and intensity of symptoms, activities with which they interfere.

- Onset of symptoms. When, where, and over what interval of time did the symptoms arise?
- *Time course of symptoms.* How did they develop? Are they constant or variable? Are there any exacerbating or alleviating factors?
- Accompanying symptoms, if present.
- Past history of similar symptoms.
- Previous illnesses and their outcome.
- Social, occupational, and family history.
- Medications, smoking, alcohol abuse, substance abuse, toxic exposures.
- Previous diagnostic studies and treatment.
- Information from third parties may be needed for patients with aphasia, confusion, dementia, or impairment of consciousness.

Physical examination. The general and neurological physical examination may yield important clues to the disease process, but only if the examiner has the requisite knowledge of the underlying principles of (neuro-)anatomy, (neuro-)physiology, and (neuro-)pathology. The examination is guided by the case history, i.e., the patient's complaints and general physical condition determine what the examiner looks for in the examination. The unselective, "shotgun" application of every possible technique of neurological examination in every patient is not only a waste of time and money; it generally only creates confusion rather than clarifying the search for the diagnosis. The neurological examination of small children, patients with personality changes or mental illness, and unconscious patients poses special challenges.

Important elements of the neurological examination include:

- Inspection. Dress, appearance, posture, movements, speech, gestures, facial expression.
- Mental Status. Orientation (to person, place, and time), attention, concentration, memory, thought processes, language function, level of consciousness.
- Cranial nerves. Olfaction, pupils, visual fields, eyegrounds, eye movements, facial movement, facial sensation, hearing, tongue movements, swallowing, speaking, reflexes.
- Motor function. Muscular atrophy/hypertrophy, spontaneous movements, coordination, paresis, tremor, dystonia, muscle tone.

- Reflexes (p. 40).
- Sensory function. The findings of sensory testing are heavily influenced by the patient's "sensitivity" and ability to cooperate. Vague sensory abnormalities without other neurological deficits are difficult to classify; their interpretation requires a good knowledge of the underlying neuroanatomy (pp. 32 ff, 106 f).
- *Posture, station, and gait.* The observation and testing of posture, station, and gait provides important information about a possible motor deficit (p. 42 ff).
- Autonomic function. The patient is questioned about bladder function, bowel movement/control, sexual function, blood pressure, cardiac function, and sweating, and is examined as needed.

Localization of the Lesion

The findings of the history and physical examination findings are then related to dysfunction of a particular neuroanatomical structure(s) (p. 2 ff) or neurophysiological process (p. 40 ff); the site of the patient's problem is thus localized (topical diagnosis).

Provision of an Etiological Diagnosis

Once the site of the problem is localized, it must be determined whether it is due to a structural lesion (e.g., hemorrhage, nerve compression, or infection) or a functional disturbance (e.g., epileptic seizure, migraine, or Parkinson disease). The line between structural and functional pathology is not perfectly defined, as there is constant interaction between these two levels; at the same time, considerations of etiology and pathogenesis also influence data interpretation. The diagnostic process ideally ends in the diagnosis of a specific disease entity (nosological diagnosis).

Additional Diagnostic Studies

The clinical diagnosis may be considered firmly established by the history and physical examination alone in many cases, e.g., migraine or Parkinson disease. Additional diagnostic studies are merely confirmatory and are generally not needed unless doubt arises as to the diagnosis, e.g., if an epileptic seizure or new type of headache should appear.

Additional studies are needed, however, if there is no other way to decide among several diagnostic possibilities remaining after thorough history-taking and physical examination. The number and type of studies needed differ from case to case. Studies that are costly or fraught with nonnegligible risk should never be ordered except to answer a clearly stated diagnostic question. The potential benefits of a proposed study must always be weighed against its risks and cost.

Laboratory Tests

Table 77, p. 407

Neurophysiological Tests

Test/Purpose	Risks	Comments
Electroencephalography: To assess elec- trical activity of the brain ¹	Surface electrodes: none Needle electrodes: infec- tion Induction of seizures by provocative methods ²	Sphenoid, subdural or depth re- cording ³ for special questions relevant to the (preoperative) diagnostic eval- uation of epilepsy
 Evoked potentials (EPs): VEPs⁴: Study of optic nerve, optic chiasm and optic tract. AEPs⁵: Study of peripheral and central segments of the auditory pathway⁶ SEPs⁷: Study of somatosensory systems⁸ MEPs⁹: Study of corticospinal motor pathway 	 None None None May induce epileptic seizures. Contraindica- tions: cardiac pacemakers, metal prostheses in the target area, pregnancy, un- stable fractures 	 Used mainly to diagnose prechiasmatic lesions Used mainly for diagnosis of multiple sclerosis, tumors of the posterior cranial fossa, brain stem lesions causing coma or brain death, and intraoperative monitoring Used to assess proximal peripheral nerve lesions (plexus, roots) and spinal cord or parietal lobe lesions Pyramidal tract lesions, motor neuron lesions, root compression, plexus, lesions, stimulation of deep nerves, differential diagnosis of psychogenic paresis
Electromyography: Study of electrical activity in muscle	Contraindication: coagulopathy. Risk of in- jury in special studies ¹¹	Provides information on motor unit disorders in patients with peripheral nerve lesions or myopathies. Not dis- ease-specific. Disposable needles should be used to prevent spread of infectious disease ¹⁰
Electroneurography: Measurement of motor and sensory conduction velocities.	Needle recordings con- traindicated in patients with coagulopathy	Localization (proximal, distal, conduc- tion block) and classification (axonal, demyelinating) of peripheral nerve le- sions ¹²
Electro-oculography: To record and assess eye movements and/or nystag- mus	Caloric testing with water contraindicated in patients with perforated eardrums	Diagnosis and localization of periph- eral and central vestibular lesions. Differentiation of saccades

1 For assessment of epilepsy, localized pathology (neoplasm, trauma, meningoencephalitis, infarct) or generalized pathology (intoxication, hypoxia, metabolic encephalopathy, Creutzfeldt-Jakob disease, coma, brain death), for sleep analysis (polysomnography), or to monitor the course of such conditions. 2 Photostimulation, hyperventilation, sleep, sleep withdrawal. 3 For diagnostic assessment before epilepsy surgery, in specialized centers. 4 Visual EPs. 5 Acoustic EPs. 6 Peripheral nerve and cochlear lesions are mainly studied audiometrically, and peripheral and central disorders by electro-oculography or posturography. 7 Somatosensory EPs. 8 Functional assessment of sensory pathways (p. 104) by tibial, median, ulnar, and trigeminal nerve stimulation. 9 Motor EPs. 10 Particularly Creutzfeldt-Jakob disease, hepatitis, AIDS. 11 *Examples*: Pneumothorax in study of the seratus anterior m., perforation of the rectal wall in study of the anal sphincter. 12 F-wave measurement for localization of proximal nerve damage, H-reflex in S1 syndrome.

Neuropsychological Tests

Comprehensive testing of cognitive function, behavior, and affective processes, perhaps in collaboration with a neuropsychologist, is required when the history and physical examination suggest the possibility of mental illness or of mental dysfunction due to neurological disease. *Objectives*: Accurate detection and effective monitoring, prognostication, identification of etiology, and treatment of mental disorders (p. 122 ff).

352

Aspect To Be Tested	Questions/Tests
• Attention (p. 116)	 Awake, somnolent, stuporous, comatose? Arousability, attention span, perception
Orientation	 Personal data (name, age, date/place of birth), orientation ("where are we?", place of residence); time (day of the week, date, month, year); situation (reason for consultation, nature of symptoms)
Memory, recall	9 The patient should be able to name the months of the year backward, spell a word backward, repeat random series of numbers between 1 and 9. Can the patient recall 3 objects mentioned 3 minutes ago, recall figures, name famous people? Tests of general knowledge
 Serial subtraction Frontal lobe function 	 Serial subtraction of 3s (or 7s), starting from 100 Perseveration¹; hand sequence test²; proverb interpretation
• Language (pp. 124, 128)	 Following commands, naming, repetition, writing, reading aloud, simple arithmetic
 Praxis Spatial orientation, visual perception 	 See p. 128 See p. 132. Naming of colors and objects

(After Schnider, 1997)

1 Drawing of simple figures (Luria's loops). 2 Command sequence: "Make a fist—open the hand to the side—open the hand flat."

Cerebrovascular Ultrasonography

Ultrasound can be used to assess the extracranial and intracranial arteries. The transmitter emits ultrasonic waves in two modes, continuous wave (CW; cross-sectional data, but no depth information) and pulse wave (PW; flow information at different levels). The reflected waves are recorded (echo impulse signal) and analyzed (frequency spectrum analysis, color coding). The flow velocity of blood particles can be determined according to the Doppler principle. As the flow velocity is correlated with the diameter of a blood vessel, its measurement reveals whether a vessel is stenotic. In direct vessel recordings. CW Doppler can be used to determine the direction of flow and the presence or absence of stenosis or occlusion. In duplex sonography, the PW Doppler and ultrasound images (echo impulse) are combined for simultaneous demonstration of blood flow (color-coded flow image) and tissue structures (tissue image). This permits visualization and quantitation of stenosis, dissection, extracranial vasculitis, and vascular anomalies. *Transcranial Doppler* (TCD) and *duplex sonography* are used to study the intracranial arteries, e.g., for stenosis, occlusion, collateral flow, vasospasm (after subarachnoid hemorrhage), shunting (arteriovenous malformation or fistula), and hemodynamic reserve.

Neuroimaging

The neuroradiologist can demonstrate structural changes associated with neurological disease with a number of different imaging techniques. When a patient is sent for a neuroimaging study, the reason for ordering the study and the question(s) to be answered by it must be clearly stated. Interventional procedures in the neuroradiology suite are mainly performed to treat vascular lesions (embolization of an arteriovenous malformation, fistula, or aneurysm; thrombolysis; angioplasty; devascularization of neoplasms; stent implantation).

Imaging Study	Indication/Objective ¹
Conventional radiography ² Skull, spine	Metallic foreign bodies, air-filled cavities, fractures, skull defects, bony anomalies, osteolysis, spinal degenerative disease
Computed tomography (CT) Head, spine, spinal canal, CT-guided diagnostic interventions, 3-D recon- struction	Assessment of skeleton (anomalies, fractures, osteolysis, degenerative changes, spinal canal stenosis), metastases, trauma, intracranial hemorrhage, cerebral ischemia, hydrocephalus, calcification, inter- vertebral disk disease, contrast studies ³ (brain, spinal canal, CT angio- graphy)
Magnetic resonance imaging (MRI) ⁴ Head, spine, spinal canal Skeletal muscle 	 Tumors (brain, spine, spinal cord), infection (encephalitis, myelitis, abscess, AIDS, multiple sclerosis), structural anomalies of the brain (epilepsy), leukodystrophy, MR angiography (aneurysm, vascular malformation), ischemia of the brain or spinal cord, spinal trauma, hydrocephalus, myelopathy, intervertebral disk disease Muscular atrophy, myositis
Angiography ^{3,5}	
Cerebral, spinal; preinterventional or preoperative study ⁶	High-grade arterial stenosis, aneurysm, arteriovenous malformation/ fistula, sinus thrombosis, vasculitis
Myelography ^{3,7}	Largely replaced by CT and, especially, MRI. Used to clarify special di- agnostic questions in spinal lesions
Diagnostic nuclear medicine Skeletal scintigraphy ("bone scan") 	• Tumor metastasis, spondylodiscitis
 CSF scintigraphy Emission tomography⁸ 	 Intradural catheter function test, CSF leak Cerebral perfusion, cerebral metabolic disorders, degenerative diseases, diagnosis of epilepsy

1 Examples. 2 Plain radiographs, X-ray tomography. 3 Risks: allergy (⇒ intolerance), latent hyperthyroidism (⇒ thyrotoxicosis), thyroid carcinoma (⇒ radioiodine therapy cannot be performed for a long time afterward), renal failure, left heart failure (⇔ pulmonary edema), plasmacytoma (⇔ renal failure). 4 Gadolinium contrast agent can be used to show blood-brain barrier lesions (e.g., acute multiple sclerosis plaques). 71-weighted scans: CSF/ edema dark (hypointense), diploe/fat light (hyperintense), white matter light; gray matter dark. 72-weighted scans: CSF/ edema dark (hypointense), diploe/fat light, muscle dark, white matter dark; gray matter light. Contrain-dications: Cardiac pacemaker, mobile ferromagnetic material. 5 Contraindicated in patients with coagulopathy. 6 Endovascular or surgical therapy. 7 Rare complications: generalized epileptic seizures, meningitis, post-lumbar puncture headache; acute transverse cord syndrome possible in patients with spinal tumors. Coagulopathy is a contraindication. 8 SPECT = single-photon emission computed tomography. Pt = positron emission tomography.

Tissue Biopsy

In certain cases, the provision of a definitive diagnosis requires biopsy of nerve (usually the sural nerve, p. 391), muscle (a moderately affected muscle in myopathy, p. 399), or blood vessels (e.g., the temporal artery in suspected temporal arteritis). These biopsies can usually be carried out under local anesthesia. Spinal tumors can be biopsied under CT or MRI guidance, and brain tumors and abscesses can be biopsied with stereotactic technique.



- Supplementary tables
- Detailed information
- Outlines
- Working aids

Table 1 Cranial nerves (p. 28)

Pathway	Cranial Nerve (CN)/Nucleus	Functions
Somatosensory (afferent)	Retina Proprioceptors of extraocular mm. ¹ Proprioceptors of extraocular mm. Semilunar ganglion, proprioceptors of masticatory, tensor veli palatini, and tensor tympani muscles Proprioceptors of extraocular mm. Geniculate ganglion VIII Vestibular ganglion; spiral ganglion X Superior ganglion	Vision Proprioception ² Proprioception Sensation in face, nose, nasal cavity, oral cavity; proprioception, dura mater (pp. 6, 94) Proprioception External ear, parts of auditory canal, outer surface of eardrum (sensation) Balance/equilibrium; hearing Middle ear, auditory tube (sensation) External auditory canal/dura mater of posterior fossa (p. 5)
Visceral (afferent)	 Olfactory cells of nasal mucosa VII Geniculate ganglion IX Inferior and superior ganglia X Inferior ganglion 	Smell Taste on anterior 2/3 of tongue (chorda tympani), taste on inferior surface of soft palate (greater petrosal n.) Taste/sensation on posterior 1/3 of tongue, pharyngeal mucosa, tonsils, audi- tory tube (sensation) Abdominal cavity (sensation), epiglottis (taste)
Motor (efferent)	III Oculomotor nucleus ³ IV Trochlear nucleus V Motor nucleus of trigeminal n. VI Abducens nucleus VII Facial nucleus IX Nucleus ambiguus X Nucleus ambiguus XI Nucleus ambiguus, motor cells of anterior horn of cervical spinal cord XII Hypoglossal nucleus	Extraocular mm. (except those supplied by CN IV, VI), raise eyelid (levator palpe- brae superioris m.) Oblique eye movements (superior oblique m.) Mastication, ⁴ tensing of palate ⁵ and tym- panic membrane ⁶ Lateral eye movements (lateral rectus m.) Facial muscles, platysma, stylohyoid and digastric muscles Pharyngeal mm., stylopharyngeus m. Swallowing (pharyngeal mm.), speech (superior laryngeal nerve) Muscles of pharynx and larynx, sternoclei- domastoid m. ⁷ trapezius m. ⁸ Muscles of tongue
Visceral (efferent)	 III Parasympathetic, Edinger–Westphal nucleus VII Parasympathetic, superior salivatory nucleus IX Parasympathetic, inferior salivatory nucleus X Parasympathetic, dorsal nucleus of vagus nerve 	Pupillary constriction (sphincter pupillae m.), accommodation (ciliary m.) Secretion of mucus, tears, and saliva (sub- lingual and submandibular glands) Secretion of saliva (parotid gland) Lungs, heart, intestine to left colonic flexure (motor); glandular secretion (res- piratory tract, intestine)

Eye muscles. 2 See p. 104. 3 Nucleus. 4 Masseter, temporalis, lateral pterygoid, and medial pterygoid muscles.
 5 Tensor veli palatini m. 6 Tensor tympani m. 7 Shoulder elevation, scapular fixation, accompanying movements of cervical spine. 8 Neck flexion and extension, head rotation.

Table 2 Segment-indicating muscles (p. 32)

Segment	Segment-indicating Muscle(s)
C4	Diaphragm
C5	Rhomboids, supraspinatus, infraspinatus, deltoid
C6	Biceps brachii, brachioradialis
C7	Triceps brachii, extensor carpi radialis, pectoralis major, flexor carpi radialis, pronator teres
C8	Abductor pollicis brevis, abductor digiti quinti, flexor carpi ulnaris, flexor pollicis brevis
L3	Quadriceps femoris, iliopsoas; adductor longus, brevis et magnus
L4	Quadriceps femoris (vastus medialis m.)
L5	Extensor hallucis longus, tibialis anterior, tibialis posterior, gluteus medius
S1	Gastrocnemius, gluteus maximus

Tables 3 Types of tremor (p. 62)

Туре	Features
Physiological tremor (PT)	Normal. Discrete, usually asymptomatic tremor of unclear significance. Isometric tremor may occur, e.g., when holding a heavy object.
Exaggerated PT, toxic or drug- induced tremor	Amplitude > PT, frequency = PT. Absent at rest. Mainly PosT. ¹ Stress (anxiety, fatigue, excitement, cold). Metabolic disturbances (hyperthyroidism, hypoglycemia, pheochromocytoma). Drugs/toxins (alcohol or drug with- drawal; mercury, manganese, lithium, valproic acid, cyclosporine A, amio- darone, flunarizine, cinnarizine, tricyclic antidepressants, neuroleptics ⇒ tar- dive tremor)
Essential tremor (ET)	Classical ET: PosT > KT ² . Approx. 60% autosomal dominant, rest sporadic. Hands > head > voice > trunk. Often improved by alcohol. Orthostatic tremor: Occurs only when standing ⇒ unsteadiness, hard to stand still. Task-specific tremor
Parkinsonian tremor	RT ³ See p. 206. Postural and kinetic tremor may also be present.
Cerebellar tremor	IT ⁴ reflecting cerebellar dysfunction. Postural tremor and head/trunk tremor may be seen when the patient is standing (alcohol intoxication).
Holmes tremor (rubral, mid- brain tremor, myorhythmia)	RT + PosT + IT, mainly proximal, disabling. Associated with lesions of nigro- striatal and cerebello-thalamic pathways (multiple sclerosis, infarct)
(Poly-)neuropathic tremor	RT, PosT, or IT, predominantly either proximal or distal. 310Hz^{5}
Palatal tremor	Symptomatic (medullary lesion due to encephalitis, multiple sclerosis, brain stem infarct) or essential; clicking noise in ear
Psychogenic tremor	Migrates from one part of the body to another. Accompanied by muscle contraction (co-contraction)

1 PosT = postural tremor. 2 KT = kinetic tremor. 3 RT = resting tremor. 4 IT = intention tremor. 5 Occurs in hereditary sensorimotor neuropathy type I, chronic demyelinating polyradiculitis, paraproteinemic neuropathy, diabetic neuropathy, and uremic neuropathy.

Table 4 Midbrain syndromes (p. 71)

Anterior Midbrain Lesions (Peduncle, Weber Syndrome)

Cause. Infarct. Less commonly caused by hemorrhage, tumor (germinoma, teratoma, pineoblastoma, astrocytoma, tentorial edge meningioma, lymphoma), or multiple sclerosis.

Structure Affected	Symptoms and Signs
Intramesencephalic fibers of oculomotor n.	Ipsilateral oculomotor paralysis + parasympathetic dysfunction (pupil dilated and unreactive to light)
Pyramidal tract	Contralateral central paralysis + face (⇒ supranuclear facial palsy) + spastic- ity. Dysarthria (supranuclear hypoglossal palsy)
Substantia nigra	Rigidity (rare)

Medial Midbrain Lesions (Tegmentum, Benedikt Syndrome)

Cause. Same as in anterior lesions.

Structure Affected	Symptoms and Signs
Intramesencephalic fibers of oculomotor n.	Ipsilateral oculomotor paralysis + parasympathetic dysfunction (see above)
Medial lemniscus	Contralateral impairment of touch, position, and vibration sense
Red nucleus	Contralateral tremor (myorhythmia \Rightarrow red nucleus syndrome, Holmes tremor)
Substantia nigra	Rigidity (variable)
Superior cerebellar peduncle	Contralateral ataxia (⇔ Claude syndrome)

Dorsal Midbrain Lesions (Tectum, Parinaud Syndrome)

Cause. Tumor of third ventricle, infarct, arteriovenous malformation, multiple sclerosis, large aneurysm of posterior fossa, trauma, shunt malfunction, metabolic diseases (Wilson disease, Niemann–Pick disease), infectious diseases (Whipple disease, AIDS)

Structure Affected	Symptoms and Signs
Oculomotor nuclei	Pathological lid retraction (Collier's sign) due to overactivity of levator palpebrae superioris m. Over the course of the disease, accommodation is impaired; the pupils become moderately dilated and unreactive to light, but they do constrict on convergence (light-near dissociation)
Medial longitudinal fasciculus	Supranuclear palsy of upward conjugate gaze (vertical gaze palsy \Rightarrow the eyes move upward on passive vertical deflection of the head, but not voluntarily). Convergence nystagmus with retraction of the eyeball on upward gaze (retraction-convergence nystagmus)
Trochlear nucleus	Trochlear nerve palsy
Aqueduct (compressed)	Hydrocephalus (headache, papilledema)

Table 4 Midbrain syndromes (continued)

Top of the Basilar Artery Syndrome

Cause. Large aneurysm of the basilar tip, thromboembolism in the upper basilar territory, vasculitis, complication of angiography. (Central paralysis is not found.)

Site of Lesion	Symptoms and Signs
Midbrain	Unilateral or bilateral vertical gaze palsy; impaired convergence; retraction nystagmus. Sudden oscillations (sensation of movement of surroundings when walking or when moving head). Collier's sign. Strabismus with di- plopia. Pupils may be constricted and responsive or dilated and unrespon- sive to light.
Thalamus, parts of temporal and occipital lobes	Visual field defects (homonymous hemianopsia, cortical blindness). Variable features: Somnolence, peduncular hallucinations (dreamlike scenic hallucinations), memory impairment, disorientation, psychomotor hyperactivity

Table 5 Pontine syndromes (p. 72)

Anterior Pontine Lesions (Ventral Pons)

Cause. Basilar artery thrombosis, hemorrhage, central pontine myelinolysis, brain stem encephalitis, tumors, trauma. Arterial hypertension (lacunar infarct).

Mid Ventral Pons

Structures Affected	Symptoms and Signs
Pyramidal tract	Contralateral central paralysis sparing the face
Intrapontine fibers of trigemi- nal nerve	Ipsilateral facial hypesthesia, peripheral-type weakness of muscles of mas- tication
Middle cerebellar peduncle	Ipsilateral ataxia

Lacunar Syndromes¹

Structures Affected	Symptoms and Signs
Pyramidal tract	Contralateral central paralysis, sometimes more pronounced in legs, with or without facial involvement
Middle cerebellar peduncle	lpsilateral ataxia, which may be accompanied by dysarthria and dysphagia, depending on the site of the lesion (dysarthria—clumsy hand syndrome)

¹Similar syndromes can also occur in patients with supratentorial lacunas (internal capsule, thalamocortical pathways).

■ Locked-in Syndrome (p. 120)

Structures Affected	Symptoms and Signs
Ventral pons (corticobulbar and corti- cospinal tracts) bilaterally, abducens nu- cleus, pontine paramedian reticular for- mation, fibers of trigeminal nerve	Quadriplegia, aphonia, inability to swallow, horizontal gaze palsy (including absence of caloric response), absence of corneal re- flex (risk of corneal ulceration)

Eyelid and vertical eye movements (supranuclear oculomotor tracts), sensation, wakefulness (reticular ascending system), and spontaneous breathing remain intact.

Appendix

Table 5 Pontine syndromes (continued)

Dorsal Pontine Lesions (Pontine Tegmentum)

Cause. Same as in lesions of ventral pons.

■ Oral (Superior) Pontine Tegmentum (Raymond-Céstan Syndrome)

Structures Affected	Symptoms and Signs
Trigeminal nucleus/fibers	Ipsilateral facial hypesthesia, peripheral paralysis of muscles of mastication
Superior cerebellar peduncle	Ipsilateral ataxia, intention tremor
Medial lemniscus	Contralateral impairment of touch, position, and vibration sense
Spinothalamic tract	Contralateral loss of pain and temperature sensation
Paramedian pontine reticular formation (PPRF, "pontine gaze center")	Ipsilateral loss of conjugate movement (loss of optokinetic and vestibular nystagmus \Rightarrow PPRF lesion with intact vestibulo-ocular reflex (VOR, p. 84))
Pyramidal tract	Contralateral central paralysis sparing the face

Caudal Pontine Tegmentum

Structures Affected	Symptoms and Signs
Pyramidal tract	Contralateral central paralysis sparing the face
Nucleus/fibers of the facial n.	Ipsilateral (nuclear = peripheral) facial palsy (⇔ Millard–Gubler syndrome)
Fibers of abducens nerve	Ipsilateral abducens paralysis (\rightleftharpoons Foville syndrome, eyes drift "away from the lesion"; loss of VOR)
Central sympathetic pathway	Ipsilateral Horner syndrome
PPRF	Loss of ipsilateral conjugate movement
Medial and lateral lemniscus	Contralateral impairment of touch, position, and vibration sense
Lateral spinothalamic tract	Contralateral impairment of pain and temperature sensation

Table 6 Medullary syndromes (p. 73)

Medial Medullary Lesions

Cause. Occlusion of the anterior spinal artery or vertebral artery.

Structures Affected	Symptoms and Signs
Hypoglossal n. nucleus/fibers	Ipsilateral peripheral (nuclear) hypoglossal paralysis
Pyramidal tract	Contralateral central paralysis sparing the face (flaccid, in isolated pyramidal tract lesions)
Medial lemniscus	Contralateral impairment of touch, position, and vibration sense (pain and temperature sensation intact)
Medial longitudinal fasciculus	Upbeat nystagmus

Lateral Medullary Lesions (Dorsolateral Medullary Syndrome, Wallenberg Syndrome)

Cause. Occlusion of posterior inferior cerebellar artery (PICA) or vertebral artery. Less common causes: tumor, metastases, hemorrhage due to vascular malformations, multiple sclerosis, vertebral artery dissection (after chiropractic maneuvers), trauma, gunshot wounds, cocaine intoxication.

Site of Lesion	Symptoms and Signs
Spinal nucleus of trigeminal nerve	Ipsilateral analgesia/thermanesthesia of the face and absence of corneal re- flex with or without facial pain
Cochlear nucleus	Ipsilateral hearing loss
Nucleus ambiguus	Ipsilateral paralysis of the pharynx and larynx (hoarseness, paralysis of the soft palate), dysarthria, and dysphagia. Tongue movement remains intact
Solitary nucleus	Ageusia (impaired sense of taste)
Dorsal nucleus of vagus n.	Tachycardia and dyspnea
Inferior vestibular nucleus	Nystagmus away from the side of the lesion, tendency to fall toward the side of the lesion, nausea and vomiting
Central tegmental tract	Ipsilateral myorhythmia of the soft palate and pharynx
Central sympathetic pathway	Ipsilateral Horner syndrome
Reticular formation	Singultus
Inferior cerebellar peduncle	Ipsilateral ataxia and intention tremor
Anterior spinocerebellar tract	Ipsilateral hypotonia
Lateral spinothalamic tract	Contralateral loss of pain and temperature sensation with sparing of touch, position, and vibration sense (sensory dissociation)

Involvement of the lower pons produces diplopia. Occipital pain in Wallenberg syndrome is most commonly due to vertebral artery dissection.

Table 7	Syndromes	affecting	the facial	muscles (p. 98	3)
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Syndrome	Etiology
Hypomimia or amimia	Basal ganglia dysfunction (p. 206), depression
Blepharospasm, Meige syndrome, lid-opening apraxia, oromandibular dystonia, tics (p. 64 ff.)	Basal ganglia dysfunction
Melkersson-Rosenthal syndrome (recurrent swelling of face/lips, peripheral facial palsy, and fissured tongue)	Unknown
Heerfordt syndrome (fever, uveitis, parotitis, peripheral facial palsy)	Occasional manifestation of sarcoidosis, lymphoma. Cryptogenic
Bilateral peripheral facial paralysis	Neuroborreliosis, Guillain–Barré syndrome, Fisher syndrome, botulism
Möbius syndrome	Congenital bilateral facial palsy and cranial nerve in- volvement (bilateral: VI; unilateral: XII, IV, VIII, IX)
Synkinesis (involuntary co-movement of facial muscles, e. g., narrowing of palpebral fissure when the lips are pursed); hemifacial spasm	Faulty regeneration of CN VII after facial palsy. Nerve root compression and segmental demyelination in hemifacial spasm
Pseudobulbar palsy	Multiple bilateral supratentorial or pontine vascular lesions
Myopathic facies	Myopathic disorders (myotonic dystrophy, my- asthenia, facial-scapular-humeral muscular dystrophy)
Gustatory sweating (Frey syndrome) or lacrimation ("crocodile tears")	Faulty regeneration of the auriculotemporal/facial nerve
Progressive facial hemiatrophy	Unknown

Table 8	Neurological	Causes	of Dy	/sphagia	(p.	102)
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Symptoms and Signs	Site of Lesion	Cause
Oral phase impaired and swallow- ing reflex delayed (slightly) be- cause of paralysis	Supratentorial Unilateral	Cerebral infarct, tumor or hemorrhage
Delayed swallowing reflex, aspira- tion (especially of fluids), pro- longed oral phase (pseudobulbar palsy, akinesia, dysarthria, dys- phonia, salivation, oromandibular dystonia)	Supratentorial Bilateral	Vascular lesions (single or multiple infarcts, hemor- rhage), trauma, tumor, multiple sclerosis, encephali- tis, parkinsonism, multiple system atrophy, Alzheimer disease, Creutzfeldt–Jakob disease, hydrocephalus, dystonia (toxic/drug-induced), chorea, intoxication, cerebral palsy
Loss of swallowing reflex, im- paired pharyngeal phase, impaired cough reflex (bulbar palsy, dy- sarthria, respiratory disturbances), risk of aspiration	Brain stem, cerebellum	Vascular lesions, multiple sclerosis, tumor, trauma, amyotrophic lateral sclerosis, syringobulbia, poliomy- elitis, Arnold–Chiari malformation, central pontine myelinolysis, listerial meningitis, spinobulbar muscu- lar atrophy, spinocerebellar degeneration
Weakness of muscles of mastica- tion, impaired oral phase, im- paired lip closure, nasal drip; im- paired pharyngeal phase (dy- sarthria) may occur: depending on which nerve/muscle is affected	Cranial nerves	Facial paralysis, Guillain–Barré syndrome, diabetic neuropathy, amyloidosis, base of skull syndrome (p. 74)
Same as above (generalized my- opathy, dysphonia)	Neuromuscular	Myasthenia, amyotrophic lateral sclerosis, Lambert– Eaton syndrome, botulism, polymyositis/dermatomy- ositis, scleroderma, hyperthyroidism, oculopharyn- geal muscular dystrophy, myotonic dystrophy, facial- scapular-humeral muscular dystrophy, nemaline my- opathy, inclusion-body myositis

Table 9 Classification of pain^{1,2} (p. 108)

Туре	Clinical Features	Etiology (examples)
Nociceptive pain (somatic, p. 110) ³	Paresthesia, allodynia, ⁴ loss of sensa- tion, readily localizable	Meralgia paresthetica, carpal tunnel syndrome, skin lesion
Neuropathic pain, neural- gia (pp. 186, 318ff.)	Severe pain in nerve distribution, paresthesiae, allodynia, sensory loss, pain on nerve pressure, readily localiz- able	Mononeuritis, polyneuropathy, trauma, nerve compression, trigemi- nal neuralgia, neuroma
Radicular pain (p. 319 f)	Same as above + aggravated by stretching (e.g., Lasègue sign) or movement	Herniated intervertebral disk, poly- radiculitis, leptomeningeal metastases, neurofibroma/schwan- noma
Referred pain	See p. 110	See p. 110
Deafferentation pain, anesthesia dolorosa	Pain in an anesthetic or analgesic nerve territory	Plexus lesion, radicular lesion, trigemi- nal nerve lesion
Phantom limb pain	Pain felt in an amputated limb	Limb amputation
Central pain	Burning, piercing pain in the region of a neurological deficit; imprecisely lo- calizable; frequently accompanied by sensory dissociation, dysesthesia, paresthesia; triggered by stimuli	Cerebral infarct, hemorrhage, or tumor (cortex, thalamus, white mat- ter, internal capsule), brain stem, spi- nal cord; syrinx, trauma, multiple sclerosis (brain stem, spinal cord)
Chronic pain ("pain disease")	Pain that lasts >6 months, impairs social contacts, emotional state, and physical activity	Sensitization of nociceptors? Transsy- naptic neuropeptide induction (calci- tonin gene-related peptide = CGRP, substance P = SP, neurokinin A = NKA)?
Psychogenic pain	Discrepancy between symptoms and organ findings and/or syndrome classification	Mental illness

1 Selected types. 2 Features may overlap. 3 = nociceptor pain. 4 Pain evoked by a normally nonpainful stimulus.

Table 10 Sleep Characteristics Observed in Sleep Studies (p. 112)

Stage	EEG ¹	EOG ²	EMG ³
Awake	α activity (8–13 Hz)	Blinks, saccades	High muscle tone, move- ment artifact
NREM stage 1	Increasing θ activity (2–7 Hz), vertex waves ⁴	Slow eye movements ⁵	Slight decrease in muscle tone
NREM stage 2	θ activity, sleep spindles, ⁶ K-complexes ⁷	No eye movement until stage 4, EEG artifact	Further decrease in muscle tone until stage 4
NREM stage 3	Groups of high-amplitude δ waves (0.5–2 Hz, amplitude > 0.75 μ V)		
NREM stage 4	Groups of high-amplitude δ waves		
REM sleep	θ activity, saw-tooth waves ⁸ may be observed	Conjugate, rapid eye move- ments (episodic)	Low to medium muscle tone

(Berger, 1992)

1 Electroencephalogram (EEG). 2 Electro-oculogram (EOG). 3 Electromyogram (EMG). 4 Steep parasagittal waves of not more than 200 μV. 5 Slow eye movements (SEM). 6 Fusiform 11.5–14 Hz waves lasting 0.5–1.5 seconds and occurring 3–8 times/second. 7 Biphasic, initially negative δ waves appearing spontaneously or in response to acoustic stimuli. 8 Grouped, regular θ activity with a saw-toothed appearance.

363

Appendix

 Table 11
 Diagnostic criteria for death ("brain death") (recommendations of the Scientific Advisory Council of the Federal Chamber of Physicians (Germany), 1998, p. 120)

Prerequisites	Acute, severe brain damage (either primary or secondary) Exclusion of other causes ¹
Clinical criteria	Coma Absent light reflex, moderately to maximally dilated pupils ² Absent oculocephalic reflex ³ Absent corneal reflex ³ Absent response to painful stimulation in trigeminal distribution Absent pharyngeal and tracheal reflexes ³ Absence of spontaneous breathing ⁴
Proof of irreversible brain damage	Required time of observation ⁵ Supratentorial primary brain damage Adults and children over 2 years of age \Rightarrow at least 12 hours Children under 2 years of age \Rightarrow at least 24 hours ⁶ Neonates \Rightarrow at least 72 hours ⁶ Infratentorial primary brain damage Same as supratentorial damage, but with at least 1 additional examination ⁶ Secondary brain damage Adults and children over 2 years of age \Rightarrow at least 72 hours Supplementary criteria ⁷ Isoelectric EEG Absence of evoked potentials ⁸ Absence of cerebral blood flow

1 Intoxication, pharmacological sedation, neuromuscular blockade, primary hypothermia, circulatory shock or coma secondary to endocrine, metabolic or infectious disease. 2 Not due to mydriatic agents. **3** See p. 26. **4** As shown by apnea test. **5** The clinical findings at the beginning and end of the observation period must be identical. **6** At least one of the following supplementary criteria must be observed twice: isoelectric EEG, loss of early acoustic evoked potentials, demonstration of absent cerebral blood flow (by Doppler ultrasound or perfusion scintigram); standardized examination procedures must be strictly followed. **7** Once the prerequisites and clinical has been met. **8** Early auditory, somatosensory cerebral or high cervical components of evoked potentials.

Table 11 a Apnea test

Steps	Measures/Objective	Parameters To Be Measured
Prerequisites	Body core temperature ≥ $36.5 ^{\circ}C^1$ Systolic blood pressure ≥ 90mmHg^2 Positive fluid balance for more than 6 hours	
Preparation	Oxygenation: Inspiratory O ₂ concentration 100% Tidal volume: 10 ml/kg body weight	PO₂ ≥200 (to 400) mmHg (54 kPa)³ pCO₂ ≤40 mmHg (5.3 kPa)
Procedure ⁴	Disconnect ventilator \Rightarrow administer 100% O ₂ at rate of 6 to 8 l/min via thin catheter (tip at carina)	Monitor heart rate, blood pres- sure, SpO ₂ , respiratory rate; ob- serve for chest or abdominal movement; check ABG every 2–3 minutes
Termination	If no respiratory activity/movement is observed in 8 minutes ⁵	$pCO_2 \ge 60 \text{ mmHg} (8 \text{ kPa}) \text{ or } pCO_2$ rises by more than 20 mmHg ⁶ (2.7 kPa)

(Wijdicks, 2001)

1 Because hypothermia impairs CO₂ production and O₂ release from oxyhemoglobin. **2** Raise with 5% albumin solution or increase intravenous dopamine when administered, if necessary. **3** Arterial blood gas (ABG) analysis. **4** If blood pressure <90 mmHg, O₂ saturation <80%, and there is severe cardiac arrhythmia, stop the apnea test and put patient back on ventilator. **5** Reconnect ventilator, ventilate at 10/min. **6** If pCO₂ baseline value of \leq 40 mmHg cannot be achieved (e.g., in patient with pulmonary disease).

Site of Lesion	Manifestations	Causes ¹
Periphery ²	Slurred speech (impaired labial/lingual articulation, rhinolalia = "speaking through nose," unclear differ- entiation of vowels and consonants), dyspnea, whis- pering (recurrent laryngeal nerve paralysis), hoarse- ness (laryngitis, vocal cord polyp, extubation)	Facial nerve palsy, myasthenia, amyotrophic lateral sclerosis, diphtheria, Guillain–Barré syn- drome, syringobulbia, tumor
Cerebellum/ brain stem (pp. 54, 70)	Ataxic dysarthria (clipped, scanning speech), articula- tion problems. Hoarse, deep voice (vagus nerve le- sion)	See p. 278 f, multiple sclerosis, in- farction
Basal ganglia (nonpyramidal dysarthria)	Hypophonia (p. 206, monotonous, soft, slurred). Spasmodic dysphonia (p. 64). Hyperkinetic speech (p. 66; explosive, loud, uncoordinated, clipped speech)	Parkinsonism, dystonia, chorea, tic, myoclonus
White matter/ cortex	Monotonous, slow, hoarse, pressured speech. Deep, variable pitch. Poor articulation	Bilateral white-matter lesions (pseudobulbar palsy, lacunar in- farcts, multiple sclerosis), uni- lateral infarct
Diffuse	Slurred, effortful, slow speech	Intoxication, metabolic distur- bances

 Table 12
 Sites and manifestations of lesions causing dysarthria (p. 130)

1 Examples; see also p. 64. 2 Pontine (bulbar paralysis), nuclear (2nd motor neuron), peripheral nerve or muscle lesion.

Type of Amnesia	Manifestations	Causes ¹ /Site of Lesion
Transient global amnesia ²	Acute onset, limited duration. Patient repeats questions (e.g., "What am I doing here?"), is helpless, anxious; can perform everyday activi- ties. Anterograde/retrograde amnesia	Ischemia (venous)? Migraine? Re- solves completely or nearly so
Acute transient amnesia	Anterograde/retrograde amnesia; manifestations of underlying disease	Complex partial seizures (focal epilepsy). Posttraumatic phenomenon
Acute persistent amnesia	Anterograde/retrograde amnesia; manifestations of underlying disease	Bilateral infarction (hippocampus, thalamus, anterior cerebral artery). Trauma (orbitofrontal, mediobasal, diencephalic). Hy- poxia (cardiopulmonary arrest, carbon monoxide poisoning)
Subacute per- sistent amnesia	Many patients are initially confused, with antero- grade/retrograde amnesia; manifestations of un- derlying disease	Wernicke–Korsakoff syndrome. Herpes simplex encephalitis. Basilar meningitis (tuberculosis, sarcoidosis, fungi)
Chronic progres- sive amnesia	Anterograde amnesia; retrograde amnesia develops in the course of the condition; manife- stations of underlying disease	Tumor (3rd ventricle, temporal lobe). Paraneoplastic "limbic" en- cephalitis (lung cancer). Alzheimer disease. Pick disease

1 Examples. 2 Also called amnestic episode.

Cause	Diagnostic criteria ¹
Alzheimer disease (p. 297 f)	Mainly temporal and parietal lobe degeneration. Rarely hereditary (autosomal dominant). No focal neurological deficit. CSF: increased acetylcholine, A β -amyloid, and τ protein levels. Cognitive deficits: Anterograde amnesia, amnestic aphasia, acalculia, impaired visuospatial performance. Behavioral changes: Hardly any at first; later anosognosia or dissimulation, paranoia, disturbance of sleep–wake cycle.
Vascular dementia (p. 298)	Subcortical vascular demyelination due to multiple infarcts, lacunes, vasculitis or CADASIL. Fluctuating course. Focal neurological signs: Hemiparesis, aphasia, apraxia, agait impairment, bladder dysfunction, Babinski sign. Pseudobulbar palsy (bilateral lesion of the corticobulbar tracts \Rightarrow dysarthrophonia or anarthria, dysphagia, lingual/facial paralysis, loss of emotional control with outbursts of laughing or crying). Cognitive deficits: Memory deficit ("forgetting to remember"), frontal brain dysfunction (p. 122). Behavioral changes: Sluggishness, reduced drive, disturbance of sleep-wake cycle
Depression	Psychomotor slowing, reduced drive, and anxiety suggest dementia, which is not, in fact, present (<i>pseudodementia</i>). See Table 42, p. 383
Alcohol	Korsakoff syndrome: Disorientation/amnesia, confabulation
Hydrocephalus	Normal pressure hydrocephalus (p. 160)
Metabolic/endocrine disorders	Wilson disease, hypothyroidism, hypopituitarism, hepatic/uremic encephalopathy, hypoglycemia, vitamin B_{12} deficiency, Wernicke encephalopathy, pellagra, hypoxia, hypoparathyroidism or hyperparathyroidism, adrenocortical insufficiency, Cushing syndrome, acute intermittent porphyria (p. 332)
Tumor	See p. 254 ff
Degenerative diseases	Parkinson disease (p. 206 ff), atypical parkinsonism (p. 302), Hunting- ton disease (p. 300), frontotemporal dementia (p. 298), hereditary ataxia (p. 280), motor neuron disease (p. 304), multiple sclerosis
Infectious diseases (p. 222 ff)	HIV and other viral encephalitides, prion diseases, neurosyphilis, Whipple disease, brain abscess, neurosarcoidosis, subacute sclerosing panencephalitis
Trauma	Chronic subdural hematoma, posttraumatic phenomenon, punch- drunk syndrome (dementia pugilistica)
Toxic	Drugs, substance abuse, heavy metal poisoning, organic toxins

Table 14 Classification of dementia (p. 136)

1 Mainly early manifestations are listed; these usually worsen and are accompanied by other manifestations as the disease progresses.

Controlled Variable(s)	Hormones	Stimulus	Effect	Comments
Water balance and blood pressure	ADH	 BP¹ Plasma osmolality 	Renal water reabsorption and arterial vasoconstric- tion (at higher ADH levels)	O ADH: Diabetes in- sipidus; OADH: ectopic production, SIADH (p. 310)
Triiodothyronine (T_3) , thyroxine (T_4)	TRH, TSH	₽ /ûT₃/T₄	TRH² increase/decrease ⇔ TSH³	DTSH (basal) usually found in primary hypothy- roidism; UTSH usually found in hyperthyroidism
Cortisol	CRH, ACTH⁴	្‡ /û Cortisol	CRH ⁵ increase/decrease	DACTH: Cushing syn- drome; DACTH: second- ary adrenocortical insuffi- ciency
Testosterone (man)	GnRH, LH, FSH⁰	口/① Te- stosterone	GnRH ⁷ increase/decrease	Testosterone: Decrease in muscle mass, loss of li- bido, hypospermia, im- potence
Estradiol, pro- gesterone (woman)	GnRH, LH, FSH	‡ /û Estradiol, pro- gesterone	GnRH increase/decrease	LH/FSH: menstrual dis- turbances, breast/uterine atrophy, osteoporosis, atherosclerosis
Prolactin (PRL)	PRL	û PRL Q PRL	û Dopamine ⁸ ↓ VIP ⁹ , TRH	ûPRL: galactorrhea, amenorrhea, headaches
Growth (somatomedins)	GHRH, GH¹⁰	Various stimuli	GHRH ¹¹ Somatostatin ¹²	Somatomedins mediate the effect of GH. DGH: acromegaly; D GH: dwar- fism (children), weight gain, muscle atrophy
Endogenous opioid peptides	β-endorphin (pituitary gland)	Various stimuli	Analgesia, food intake, thermoregulation, learn- ing, memory	

Table 15 The hypothalamic-pituitary regulatory axis (p. 142)

1 Blood pressure. **2** Thyrotropin-releasing hormone. **3** Thyroid-stimulating hormone of the anterior pituitary = thyrotropin. **4** Adrenocorticotropic hormone of the anterior pituitary = corticotropin. **5** Corticotropin-releasing hormone (bth anterior pituitary hormones); both are called *gonadotropins*. **7** Gonadotropin-releasing hormone. **8** Released from the hypothalamus; inhibits prolactin release via pituitary D₂ receptors. **9** Vasoactive intestinal peptide (anterior pituitary). **10** GH = human growth hormone. **11** Growth hormone-releasing hormone (stimulatory). **12** Released from hypothalamus (inhibitory).

Table 16 Limbic syndromes (p. 144	Table 16	Limbic	syndromes	(p.	144
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Syndrome	Symptoms and Signs	Site of Lesion ¹
Delirium, acute confusional state	Disturbances of consciousness, attention, perception, memory, sleep-wake cycle, and cognition. Visual hallucinations. Fluc- tuating motor hypoactivity and hyperac- tivity. Affective disturbances (anxiety, de- pression, irritability, euphoria, helpless- ness)	Bilateral mediobasal temporal lobe (hippocampus, amyg- dala), hypothalamus
Pathological laughing and crying	Uncontrollable emotional outbreaks. Seen in central paralysis (pseudobulbar palsy, amyotrophic lateral sclerosis, multiple sclerosis) and focal epilepsy (gelastic seizures). Stroke prodrome	Internal capsule, basal ganglia, thalamus, corticonuclear tract
Aggressive, violent behavior; fits of rage	Aggression with minimal or no provoca- tion, seen in focal epilepsy, head trauma, hypoxic encephalopathy, brain tumor, herpes simplex encephalitis, rabies, cere- bral infarction or hemorrhage, hypogly- cemia, intoxication (drugs, alcohol)	Mediobasal temporal lobe (amygdala)
Indifference, apathy, akinetic mutism	Usually due to a primary illness such as Alzheimer or Pick disease, herpes simplex or AIDS encephalitis, hypoxic en- cephalopathy, cerebral infarction or hemorrhage	Bilateral septal area, cingulate gyrus
Memory deficit, transitory global amnesia (p. 134, Table 13)	Korsakoff syndrome: impairment of short- term memory and sense of time. Other cognitive functions and consciousness are unimpaired	Both mamillary bodies, mediobasal temporal lobe
Disturbed sexuality	Hypersexual behavior: after head trauma or stroke, or as a side effect of dopamin- ergic antiparkinsonian medication. Diminished libido: depression, medica- tions	Septal area, hypothalamus

 ${\bf 1}$ The specified lesions do not always produce these syndromes, and the location and extent of the causative lesion is often not known with certainty.

Table 17 Tests for autonomic circulatory dysfunction (p. 148)

Dizziness, syncope, lack of concentration, forgetfulness or tinnitus should not be attributed to circulatory dysfunction unless thorough diagnostic evaluation reveals a circulatory cause (see p. 166 ff).

Test	Method	Normal Findings
Tests of sympathetic function		
Schellong test (orthostasis test)	Measure heart rate and blood pressure once per minute for 10 minutes with the patient supine, then 5 minutes with the patient erect (tilt table if necessary)	Heart rate rises by no more than 20 beats/min; systolic blood pressure drops no more than 10 mmHg, diastolic no more than 5 mmHg
Valsalva maneuver (VM)	The patient inspires deeply and tries to exhale against resistance (up to 40 mmHg) for 10–15 seconds. The blood pressure is measured before, during and after the VM, or continuously	During VM, the blood pressure does not drop below 50% of initial value; after VM, the blood pressure rebounds above the initial value (+ reflex bradycardia)
Hand grip test	Isometric muscle contraction (hand grip) 30% of maximal force for 3 minutes	Diastolic blood pressure > 15 mmHg
Tests of parasympathetic func- tion		
Schellong test with 30/15 quotient	Continuous ECG recording	Quotient of R-R interval of 30th and 15th heart beat after standing up is > 1
Respiratory sinus arrhythmia (respiratory test)	ECG recorded with patient supine and breathing maximally deeply at 6/min for a total of 8 cycles. Repeat after a period of rest	The quotient of the longest (expiration) and shortest R-R intervals is normally > 1.2 (age-dependent)
VM with ECG	Continuous ECG recording	Quotient of longest and short- est R-R interval is > 1 (age-de- pendent)
Carotid sinus massage ¹	Perform unilateral carotid sinus massage for 10–20 seconds while continuously monitoring blood pressure and ECG with emergency resuscitation equipment ready	Reflex decrease in heart rate and blood pressure

(Low, 1997)

1 Contraindicated in carotid stenosis. Risk of cardiac arrest and ischemic stroke.

Table 18 Respiratory disturbances in neurological disease (p. 150)

Hypoventilation	
Site of lesion	Causes
Brain stem, upper cervical spinal cord	Tumor, infarction, hemorrhage, meningoencephalitis (<i>Lis- teria</i> , poliomyelitis), trauma, multiple sclerosis, intoxica- tion, parkinsonism (rigidity of respiratory muscles)
Motor anterior horn cell	Amyotrophic lateral sclerosis, tetanus, poliomyelitis, post- polio syndrome
Peripheral nerve	Guillain-Barré syndrome, phrenic nerve lesion
Neuromuscular	Myasthenia gravis, botulism, Lambert–Eaton syndrome, muscular dystrophy, polymyositis, acid maltase deficiency, electrolyte imbalance (NaU, KU, CaU, phosphateU, MgU)
Hyperventilation	
Possible metabolic changes	
Ca^{2*} [] \Rightarrow carpopedal spasm, paresthesiae, tetany; <i>Phosphate</i> [] \Rightarrow weakness; pH (] \Rightarrow dizziness, visual disturbances, syncope, seizures	Pneumonia, pulmonary embolism, asthma, acidosis (dia- betic, renal, lactate 1), meningoencephalitis, brain tumor, fever, sepsis, salicylates, anxiety, pain, psychogenic

Table 19 Neurological causes of gastrointestinal dysfunction (p. 154)

GI Syndrome ¹	Manifestations	Causes ²
Dysphagia	See p. 102	Table 8
Gastroparesis	Delayed gastric emptying ⇔ nausea, vomiting, anorexia, bloat- ing	Diabetes mellitus, amyloidosis, paraneoplastic syn- drome, dermatomyositis, Duchenne-type muscular dystrophy
Intestinal pseudo- obstruction	Impaired intestinal motility ⇔ nausea, vomiting, bloating, weight loss, impairment of peristalsis	Parkinson disease, multiple sclerosis, transverse spi- nal cord syndrome, Guillain–Barré syndrome, dia- betes mellitus, botulism, amyloidosis, paraneoplastic syndrome, drugs (tricyclic antidepressants, codeine, morphine, clonidine, phenothiazines, anticholiner- gics, vincristine), Hirschsprung disease
Constipation ³	Highly variable. Infrequent hard stools, straining, bloating, sensa- tion of incomplete defecation, pain, flatulence, belching	Lack of exercise, dysphagia, poor nutrition, trans- verse spinal cord syndrome, head trauma, brain stem lesions, Parkinson disease, multiple system atrophy, multiple sclerosis, diabetes mellitus, porphyria, drugs (morphine, codeine, tricyclic antidepressants)
Diarrhea ⁴	¹ Defecation rate, liquid stools, tenesmus	Diabetes mellitus, amyloidosis, HIV infection, drugs, Whipple disease
Vomiting⁵	Gagging, yawning, nausea, hyper- salivation, pale skin, outbreaks of sweating, apathy, low blood pres- sure, tachycardia	Intracranial hypertension (p. 158), vertigo (p. 58), mi- graine, infectious and neoplastic meningitis, drugs (digitalis, opiates, chemotherapeutic agents), intox- ication
Anal inconti- nence	Complete or partial loss of control of defecation	Diabetes mellitus, multiple sclerosis, spinal cord le- sions, lesion of the conus medullaris or cauda equina, dementia, frontal lobe lesions (tumor, infarction)

1 Gastrointestinal syndrome. 2 Neurological diseases often associated with gastrointestinal syndromes or neurogenic causes of such syndromes. 3 Normal frequency of defecation is ca. 3 times a week (variable). 4. Upper limit of normal frequency of defecation, ca. 3 times/day. In *diarrhea*, the stool weight is >200 g/day. In *pseudodiarrhea*, the defecation rate in increased but the stool weight is not. Diarrhea must be differentiated from anal incontinence. 5 Vomiting is regulated by a "vomiting center" in the reticular formation, which lies between the olive and solitary tract. *Input*: Chemoreceptors of the area postrema (p. 140), vestibular system, cortex, limbic system, gastrointestinal and somatosensory afferents. *Output*: Phrenic nerve (diaphragm), spinal nerve roots (respiratory and abdominal musculature), vagus nerve (larynx, pharynx, esophagus, stomach).

Site of Neuro- logical Lesion	Neurological Condition	Type of Bladder Dysfunction
Supratentorial	Stroke (frontal cortex, motor pathway)	Frequency \hat{u} , urge \hat{u}^{1} , urge incontinence ² , detrusor hyperreflexia
	Parkinson disease	Detrusor hyperreflexia, bladder hypocontractility
	Frontal brain tumor	Frequency û, urge û, urge incontinence
	Dementia	Usually a late manifestation: frequency $\hat{\mathbf{u}}_{\!\!\!,}$ urge incontinence
Supratentorial and infraten- torial	Multiple sclerosis ³ (variable distur- bances depending on site of plaques)	Frequency \hat{U} , urge \hat{U} , imperative urinary urge, urge incontinence, detrusor hyperreflexia, DSD ⁴
	Amyotrophic lateral sclerosis	Frequency $\hat{\mathbf{u}}$, urge incontinence, detrusor hyperreflexia
	Multiple system atrophy	Nocturia, frequency $\hat{\mathbf{u}}_{\!\!\!\!}$ urge incontinence, impairment of voluntary voiding
Spinal cord⁵	Trauma, tumor, ischemia, myelitis, multiple sclerosis, cervical my- elopathy, spinal arteriovenous fistula	Lesion above S2 ("reflex bladder") ⇒ hyperreflexia, residual urine, DSD Lesion of sacral micturition center ("autonomic blad- der") ⇒ residual urine, detrusor areflexia, impaired bladder reflex
Cauda equina, peripheral nerves	Autonomic neuropathy (e.g., dia- betes mellitus, paraneoplastic syn- drome, Guillain-Barré syndrome, drugs, toxins), trauma, lumbar canal stenosis, myelodysplasia, tumor, herpes zoster, arachnoid- itis, disk hemiation	Residual urine, detrusor areflexia, impaired bladder reflex, impaired filling sensation, frequency $\ensuremath{\mathbb{Q}}$

Table 20 Neurogenic bladder dysfunction (p. 156)

1 Pollakisuria. 2 Urge incontinence \Rightarrow involuntary passage of urine on strong (imperative) urinary urge. 3 Urinary tract infections are frequent. 4 Detrusor-sphincter dyssynergy. 5 Spinal shock with detrusor areflexia (bladder atony, "shock bladder") and residual urine formation (overflow incontinence). Overdistention of the bladder can lead to a sharp rise in blood pressure accompanied by headache, dizziness, and hyperhidrosis above the level of the spinal lesion.

Table 21 Causes of intracranial hypertension (p. 158)

Pathogenetic Mechanism	Causes
Mass lesion	Hematoma (epidural, subdural, intracerebral), brain tumor/ metastasis, brain abscess
CSF outflow obstruction	Hydrocephalus
Increased brain volume	Pseudotumor cerebri, infarct, global hypoxia or ischemia, hepatic encephalopathy, acute hyponatremia
Increased brain volume and increased intravascular blood volume	Head trauma, meningitis, encephalitis, eclampsia, hyper- tensive encephalopathy, venous sinus thrombosis

Appendix

Pathogenetic Mechanism	Findings	Causes
Impaired spinal CSF circulation ¹	Absence of pressure rise in Queckenstedt test	Tumor, arachnoiditis, herniated intervertebral disk
CSF leak	Spinal dural defect, rhinor- rhea, otorrhea	Prior LP ² , trauma, neurosurgical procedures, tumor, osteomyelitis
Dehydration	Thirst, dry skin, fatigue, low blood pressure, light- headedness or uncon- sciousness	Isotonic, hypotonic: Vomiting, diarrhea, diuretics. Hyper- tonic: Thirst, profuse sweating, mannitol administration
Spontaneous	Low ICP	Spinal CSF fistula (?)

 Table 22
 Causes of intracranial hypotension (p. 160)

1 In a complete blockade, the pressure decreases sharply when small volumes of fluid are removed. In that case, there is a risk of aggravation of spinal compression syndrome due to incarceration. 2 Lumbar puncture.

 Table 22 a
 Causes of transient monocular blindness (amaurosis fugax, p. 168)
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Site and Type of Lesion	Cause
Retinal vessels—embolic	Atheroembolic/thromboembolic (e.g., internal carotid artery dissection/stenosis), cardioembolic (right–left shunt, e.g., in patent foramen ovale, thrombus in atrial fibrillation, mitral valve defect, acute myocardial infarction, endocarditis, artificial heart valve)
Retinal vessels—ischemic	Low perfusion pressure (orthostatic hypotension, arteriovenous shunt, intracranial hypertension, glaucoma); high perfusion resistance (migraine, glaucoma, malig- nant arterial hypertension, increased blood viscosity, retinal venous thrombosis, vasospasm)
Retina	Retinal detachment, paraneoplastic (p. 388), chorioretinitis, blow to eye
Orbit/eyeball	Tumor, subluxation of lens, vitreous body hemorrhage
Optic nerve	Vascular (ischemia, arteritis [p. 180], malignant arterial hypertension), papil- ledema, retrobulbar neuritis (Uhthoff sign, p. 216)
Unknown	Blowing the nose, malaria, pregnancy, hypersensitivity to cold, interleukin-2, acute stabbing pain, sinus lavage

(Gautier, 1993; Warlow et al., 2001)

Table 23	Causes	of chronic	daily	headache	(p. 182)	
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Type of Headache	Symptoms and Signs/Syndromes
Primary headache	
Chronic tension headache	See p. 182
Migraine	Mild to severe, often unilateral pain (transformed migraine). Additional migraine at- tacks (p. 184)
Atypical facial pain	Unilateral or bilateral pain, often predominantly felt in the nasolabial or palatal region, often very severe. Unresponsive to a wide variety of medical and surgical therapies. Normal findings on a wide variety of diagnostic tests ("diagnosis of exclusion")
Secondary headache	Posttraumatic, drug-induced, vascular (p. 182), intracranial mass, hydrocephalus, sinusi- tis, parkinsonism, cervical dystonia, myoarthropathy of the masticatory apparatus, 1 mental illness (depression, schizophrenia, hypochondria), cervical spine lesions (degenerative lesions, fractures, Klippel-Feil syndrome), Down syndrome, basilar im- pression, osteoporosis, skull metastasis, spondylitis, nheumatoid arthritis, lesions of cer- vical spinal cord/meningismus (tumor, hemorrhage, syringomyelia, cervical myelopathy, von Hippel-Lindau syndrome, meningitis, carcinomatous meningitis, intracranial hypo- tension)

1 Temporomandibular joint dysfunction, oromandibular dysfunction.

Table 24 Prognostic factors in epilepsy (p. 198)

Favorable Prognostic Factors	Unfavorable Prognostic Factors
One seizure type	Multiple seizure types
No interictal neurological deficit	Interictal neurological deficit
Older age of onset	Younger age of onset
Seizures secondary to a treatable disease	Spontaneous seizures
Individual seizures of short duration	Status epilepticus
Frequent seizures	Infrequent seizures
Good response to anticonvulsants	Poor response to anticonvulsants

(Neville, 1997)

Table 25 Causes of syncope (p. 200)

Cause	Underlying Condition/Trigger
Cardiac	Arrhythmia (bradyrhythmia, tachyrhythmia, or reflex arrhythmia), heart disease (e.g., cardiomyopathy, myxoma, mitral stenosis, congenital malformation, pulmonary embolism)
Hemodynamic	Hypovolemia, hypotension (vasovagal as an emotional reaction to pain, anxiety, sudden shock, sight of blood; hypotension from prolonged standing, heat, exhaustion, alcohol; multiple system atrophy; polyneuropathies, e.g., amyloid, hereditary, toxic; polyradicu- lar neuropathies/Guillain-Barré syndrome; antihypertensive agents, nitrates, other drugs; postural orthostatic tachycardia syndrome = POTS; paraplegia above T6)
Cerebrovascular	Subclavian steal syndrome, basilar migraine, Takayasu disease
Metabolic	Hypoglycemia, hyperventilation, anemia, anoxia, postprandial (older individuals)
Miscellaneous	Coughing fit (cough syncope = tussive syncope = laryngeal syncope), micturition (mic- turition syncope), defecation, prolonged laughing ("laughing fit", geloplegia), glos- sopharyngeal neuralgia, affect-induced respiratory convulsion in childhood (breath- holding spells), pop concerts (teenage females), lying in supine position during preg- nancy (supine syndrome)

(Bruni, 1996; Lempert, 1997)

Table 26 Causes of sudden falling without loss of consciousness (p. 204)

Type of Fall/Pathogenesis	Cause	Features
Drop attack	TIA ¹ in vertebrobasilar territory	Usually accompanied by dizziness, diplopia, ataxia, or paresthesias
	TIA in anterior cerebral artery territory	Seen when the two anterior cere- bral arteries arise from a common trunk
	Colloid cyst of 3rd ventricle	Position-dependent headache
	Posterior fossa tumor	Sudden fall after flexion of neck
Parkinsonism	Parkinson disease, multiple system atrophy	See p. 206 f
Muscle weakness	Myopathy, Guillain–Barré syn- drome, polyneuropathy, spinal lesions	See p. 50 f
Spinal or cerebellar ataxia, gait apraxia	Funicular myelosis, cerebellar lesions, metabolic encephalo- pathies, hydrocephalus, lacunar state, cervical myelopathy, multi- ple sclerosis	See specific diseases
Cryptogenic	Unknown	Occurs in women over 40 while walking
Vestibular disorder	Ménière disease (vestibular drop attack ⇔ Tumarkin otolithic crisis); occasionally due to otitis media, toxic or traumatic causes	Dizziness, nausea, nystagmus, tin- nitus. Vestibular drop attacks may occur in isolation
Cataplexy	Loss of muscle tone triggered by emotional stimuli (fright, laughter, anger)	Alone or with narcolepsy

1 Transient ischemic attack.

 Table 27
 Diagnostic criteria for multiple sclerosis (pp. 216, 218)

Manifestations	Additional Information Needed for Diagnosis
Two or more episodes; objective eviden- ce ¹ of 2 or more lesions	None
Two or more episodes; objective evidence of 1 lesion	Disseminated lesions (MRI ²) or two or more MS-typical lesions (MRI and positive CSF tests ³) or relapse ⁴
One episode; objective evidence of 2 or more lesions	Dissemination of lesions over time (MRI ^s) or relapse
One episode; objective evidence of 1 le- sion (monosymptomatic syndrome)	Disseminated lesions (MRI ²) or two or more MS-typical lesions (MRI + positive CSF findings ³) + dissemination of lesions over time (MRI ⁵) or relapse
Gradual worsening of neurological manifestations suggestive of MS	Positive CSF findings ³ + disseminated lesions ⁶ or pathological VEP + 4–8 cerebral lesions ⁷ + dissemination of lesions over time (MRI ⁵) or continuous progression for 1 year

(McDonald et al., 2001)

1 MRI, CSF, visual evoked potentials (VEP). 2 For special criteria, see McDonald et al., 2001. 3 Oligoclonal immunoglobulin, elevated IgG index. 4 Topographic-anatomic classification differs from that of previous episodes. 5 Follow-up examination after an interval of at least 3 months; for special criteria, see McDonald et al., 2001. 6 Nine or more cerebral lesions or two or more spinal lesions or 4–8 cerebral lesions. 7 In patients with fewer than 4 cerebral lesions, at least 1 additional spinal lesion must be observed by MRI.

Clinical Features	Additional Findings	Treatment ¹
Previously healthy patient	Gram-positive cocci in CSF	Vancomycin + cephalosporin
	Gram-negative cocci in CSF	Penicillin G
	Gram-positive bacilli in CSF	Ampicillin or penicillin G + aminoglycoside ³
	Gram-negative bacilli in CSF	Cephalosporin ² + aminoglycoside ³
Previously healthy patient	Negative CSF Gram stain ⁵	Bacterial infection suspected: cephalosporin ² . Age > 50 years + ampicillin Viral infection suspected: influenza A, amantadine or rimantadine; herpes simplex (p. 236); cytome- galovirus (p. 244); poliovirus (p. 242); HIV (p. 241); varicella zoster (p. 239)
Septic focus (e. g., mastoiditis), neurosurgery, head trauma	Supportive evidence from imaging study, e.g., CT with bone windows	Vancomycin + cephalosporin ²
Immune deficiency/immunosup- pressant therapy ⁴	(possible) Brain stem signs, Gram- negative bacilli in CSF	Ampicillin + ceftazidime
Nosocomial infection	(possible) Gram-negative bacilli in CSF	Cephalosporin ² + e.g., oxacillin or fosfomycin + aminoglycoside ³
Focal neurological signs	Temporal lobe process demon- strated by EEG, CT and/or MRI	Acyclovir (p. 236)
Spinal and radicular pain	Spinal epidural abscess confirmed by imaging study (MRI, myelogra- phy, or CT)	Cephalosporin ² + (e.g.) oxacillin or fosfomycin + aminoglycoside ³ . Surgery
Neonate (<3 months of age)	Negative CSF Gram stain	Ampicillin + cephalosporin ²

Table 28 Therapeutic guidelines for meningoencephalitis (p. 224)

1 Drugs recommended by Quagliarello and Scheld (1997). **2** E.g., cefotaxime or ceftriaxone. **3** Gentamycin or tobramycin. **4** Predisposes to tubercular, fungal, and other opportunistic infections (pp. 233, 245 f). **5** Aseptic/viral meningitis, p. 234.

Pathogen	Portal of Entry/Focus	Clinical Features
Pneumococcus (S. pneumoniae/ Gram ⁺ extracellular diplococ- cus) ⇔ adults ¹	Nasal and pharyngeal mucosa, head trauma, neurosurgical procedures, external CSF drainage	Meningitis may be accompanied or preceded by sinusitis, otitis media or pneumonia. Posttraumatic meningitis may occur several years after trauma; re- current meningitis (CSF leak? Im- munodeficiency?). Course may be hyper- acute (nonpurulent meningitis?), acute or subacute (days to weeks). Epileptic seizures. Risk of brain abscess, subdural empyema or cerebral vasculitis
Meningococcus (N. meningitid- is³/Gram- intracellular diplo- coccus) ⇒ children and ado- lescents ⁴	Nasopharynx	Hyperacute course with sepsis, adrenocortical insufficiency and consump- tion coagulopathy (Waterhouse-Frider- ichsen syndrome). Petechial or confluent cutaneous hemorrhages. Myocarditis/peri- carditis
Haemophilus influenzae (Gram- bacillus) ⇔ children and ado- lescents	Nasal and pharyngeal mucosa	Usually type B. May be accompanied or preceded by sinusitis, otitis media, or pneumonia
Listeria (L. monocytogenes/ Gram¹/organism difficult to identify) ⇔ neonates ⁵ , adults > 50 years of age	Gastrointestinal tract (con- taminated food, e.g., dairy products or salads)	Focal neurological deficits, particularly brain stem encephalitis (rhombencephali- tis), are commonly seen. Predisposing factors: pregnancy, old age, alcoholism, immune suppression, primary malignancy. CSF findings are extremely variable ("mixed cell picture")
Staphylococcus (S. aureus/ Gram ⁺) ⇒ neonates and adults Enterobacter (Gram-/bacilli) ⇒ neonates	Endocarditis, head trauma, ex- ternal CSF drainage, lumbar puncture, urinary tract, spondylodiskitis	In association with sepsis, i. v. drug use, alcoholism, diabetes mellitus, primary malignancy
<i>M. tuberculosis</i> (acid-fast bacil- lus)	Extracerebral organ tuberculo- sis	See p. 232

Table 29 Bacteria commonly causing meningitis and meningoencephalitis (p. 226)

Gram⁺ = Gram-positive; Gram⁻ = Gram-negative.

1 Åge > 18 years. 2 Very rapidly progressive meningitis, low cell count and high total protein and lactate levels in CSF, and CSF smear culture containing large quantities of bacteria. 3 Group A, Central Africa, South America; Group B, Europe; Group C, North America; type may change. 4 Age 3 months to 18 years. 5 Age ≤ 3 months.

Table 30	Viruses	causing	CNS	infection	(p.	234))
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Common	Occasional	Rare
Meningitis		
Enteroviruses ^{1,3} , arboviruses ^{2,3} , HIV ⁵ , HSV type 2 ⁴	HSV type 14, LCMV3, mumps virus ³	Adenoviruses ⁴ , CMV ⁴ , Epstein–Barr virus (EBV) ⁴ , influenza virus A+B ³ , measles virus ³ , parainfluenza virus ³ , rubella virus ³ , varicella-zoster virus (VZV) ⁴
Encephalitis, myelitis		
Arboviruses, enteroviruses, HSV type 1, mumps virus	CMV, EBV, HIV, measles virus ³ , VZV	Adenoviruses, influenza A, LCMV, parain- fluenza virus, rabies virus ³ , rubella virus, HTLV-I ^{5,6}

1 Poliovirus 1–3, coxsackievirus (B5, A9, B3, B4, B1, B6), echovirus (7, 9, 11 30, 4, 6, 18, 2, 3, 12, 22), enterovirus (70, 71). **2** Arthropod-borne viruses, including alphaviruses, flaviviruses, pestiviruses, bunyaviruses, and orbiviruses. **3** RNA virus. **4** DNA virus. **5** Retrovirus. **6** Human T-cell lymphotropic virus type I causes myelitis.

 Table 31
 Grades of malignancy of brain tumors (p. 264)

Tumor	Grade I (benign)	Grade II (semi- benign)	Grade III (malig- nant)	Grade IV (malig- nant)
Neuroepithelial tumors Astrocytoma Fibrillary, protoplasmic, gemistocytic astrocytoma Anaplastic astrocytoma Glioblastoma (= glioblastoma multiforme) Pilocytic astrocytoma Pleomorphic xanthoastrocytoma	+++	+++ +++	++ +++ +	+++
Oligodendroglioma • Oligodendroglioma • Anaplastic oligodendroglioma		+++	+++	
Ependymoma • Ependymoma (cellular, papillary, epithelial) • Anaplastic ependymoma	++	+++	++ +++	
Mixed glioma • Oligoastrocytoma • Anaplastic oligoastrocytoma		+++	+++	
Choroid plexus tumors Plexus papilloma Plexus carcinoma	+++		+++	++
Neuronal/mixed neuronal-glial tumors • Gangliocytoma • Ganglioglioma • Anaplastic ganglioglioma	+++	+++	+++	
Pineal tumors • Pineocytoma • Pineoblastoma (PNET)	+++	++		+++
Embryonal tumors • Primitive neuroectodermal tumor (PNET), see p. 260 • Neuroblastoma				+++ +++
Cranial nerve tumors • Schwannoma	+++			
Meningeal tumors Meningioma Anaplastic meningioma	+++		+++	
Blood vessel tumors • Hemangiopericytoma • Glomus tumor	+++	+++	++	
Lymphoma • Primary CNS lymphoma			++	+++
Germ cell tumors • Germinoma			+++	
Intra- and suprasellar tumors • Pituitary adenoma • Craniopharyngioma	+++ +++		+	
Metastatic tumors				+++

(Kleihues et al., 1993 and Krauseneck, 1997)

+++, Common; ++, Rare; +, Very rare.

Table 32 Karnofsky performance scale for quantification of disability (p. 264)

General Condition	%	Comments
Patient can perform normal daily activities and work without impairment	100	Normal; no complaints; no evidence of disease
No specific treatment required	90	Able to carry on normal activity; minor impair- ment
	80	Normal activity with effort; some impairment is clearly evident
Patient cannot work; can meet most personal needs, but needs some degree of assistance; can be cared for at home	70	Cares for self; cannot perform normal activities or work
	60	Needs occasional assistance, but can meet most personal needs
	50	Needs considerable assistance and frequent medical care
Patient cannot care for self; needs to be cared for in a hospital, nursing home, or at home by a nurse/family members. Disease may progress rapidly	40	Disabled; requires special care and assistance; home nursing care still possible
Taplay	30	Severely disabled; hospitalization indicated al-
	20 10	Gravely ill; hospitalization necessary Moribund
in a hospital, nursing home, or at home by a nurse/family members. Disease may progress rapidly	20	Severely disabled; hospitalization indicated al- though death not imminent Gravely ill; hospitalization necessary

Table 33 Glasgow coma scale (p. 266)

l Eye Opening	Score	ll Best Verbal Response	Score	III Best Motor Response (arms)	Score
				Obeys commands	6
		Oriented	5	Selectively avoids painful stimuli	5
Spontaneous	4	Confused	4	Withdraws limb from painful stimuli	4
To speech	3	Single words	3	Flexes limb in response to painful stimuli	3
To pain	2	Meaningless utterances	2	Extends limb in response to pain- ful stimuli	2
No response	1	No response	1	No response	1

(Teasdale, 1995)

The scores in columns I, II, and III are summed to yield the overall value. GCS 13–15 = mild head trauma; GCS 9–12 = moderate head trauma; GCS 3–8 = severe head trauma.

Table 34	Criteria for	assessment	of head	trauma	(p. 266)	
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Severity (GCS) ¹	Risk of Secondary Injury ²	Symptoms and Signs ³
Mild (13–15)	Low	Impairment of consciousness lasting < 1 hour Asymptomatic (or, at most: headache, dizziness, bruises, and lacera- tions)
Moderate (9–12)	Moderate	Impairment of consciousness at time of accident or thereafter, last- ing between 1 and 24 hours Increasingly severe headache Alcohol/drug intoxication No reliable description of accident. Multiple trauma, severe facial injur- ies, basilar skull fracture, suspicion of depressed skull fracture or open head injury. Posttraumatic ⇒ epileptic seizure, vomiting, annesia Age < 2 years (except in minor accidents), possibility of child abuse
Severe (3–8)	High	Impairment of consciousness lasting < 24 hours + brain stem syn- drome or Impairment of consciousness lasting > 24 hours or Posttraumatic psychosis lasting > 24 hours Impairment of consciousness not due to alcohol/substance abuse/medi- cations, and not a postictal or metabolic phenomenon Focal neurological signs Depressed skull fracture, open head injury

(White and Likavec, 1992) 1 Glasgow Coma Scale. 2 Patients with one or more manifestations from the list at right belong to the corre-sponding risk group. 3 Criteria for assessment of severity are in **bold type**.

Table 35	Late complications	of head	trauma (p	o. 268)
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Complications	Clinical Features	Remarks
Posttraumatic syn- drome	Headache, nausea, vertigo, orthostatic hypotension, depressed mood, irritability, fatigue, insomnia, impaired concentration	Usually follows mild head trauma; may cause significant psychosocial impairment
Chronic subdural hematoma (SDH)	Headache, behavioral change, focal signs	Usually follows mild trauma (predisposing factors: old age, brain atrophy, alco- holism)
Subdural hygroma	Same as in chronic SDH	Symptoms may improve when the patient is lying down and worsen on standing
CSF leak	Drainage of CSF from the nose or ear; risk of recurrent meningitis, brain abscess	CSF rhinorrhea worsens on head flexion. CSF otorrhea indicates a laterobasal skull fracture
Hydrocephalus	Headache, behavioral change, urinary in- continence	Normal pressure hydrocephalus, venous sinus thrombosis
Epilepsy	Focal/generalized seizures	May arise years after head trauma
Encephalopathy	Behavioral changes	See p. 122 ff. Types include septic en- cephalopathy, punch-drunk en- cephalopathy (p. 302, Table 44)
Critical illness neu- ropathy and my- opathy	Prolonged ventilator dependence, weak- ness	Associated with sepsis and multiple organ failure
Heterotopic ossifi- cation (myositis ossificans)	Restricted mobility of joints, pain	Due to muscle trauma
Complications of immobility	Bed sores, peripheral nerve lesion, joint malposition	Ensure proper positioning and frequent changes of position (especially of para- lyzed limbs)

379

Table 36 Spinal fractures (p. 272)

Fracture/Dislocation	Pathogenesis	Stability ¹
Cervical spine		
 Atlantoaxial dislocation Jefferson's fracture² Dens fracture Bilateral axis arch fracture⁴ Dislocation fracture of C3-7 Lateral compression fracture 	 Dislocation between C1 and C2 Axial trauma Hyperflexion Hyperflexion and distraction Hyperflexion Flexion and axial compression 	 Unstable Unstable³ Unstable Unstable Stable
Thoracic spine, lumbar spine		
Compression fracture	Fall (back, buttocks, extended legs), direct trauma. These fractures may be pathological (osteoporosis, myeloma, metastasis)	• Stable
Burst fractureDislocation fracture		StableUnstable

(Ogilvy and Heros, 1993; Sartor 2001)

1 At the time of injury. 2 Fracture of the ring of C1 due to compression between the occiput and C2. 3 May be overlooked if the dens is not displaced; sometimes stable. 4 Hangman's fracture.

Table 37 Classification of traumatic transverse spinal cord syndrome (p. 274)

Loss of Function	Category	Features
Complete	А	No sensory or motor function, including in S4–5
Incomplete	В	No motor function. Sensory function intact below level of lesion, including in S4–5
Incomplete	С	There is motor function below level of lesion; most segment-in-dicating muscles have strength ${<}3$
Incomplete	D	There is motor function below level of lesion; most segment-indicating muscles have strength ≥ 3
None	E	Normal motor and sensory function

(American Spinal Cord Injury Association Impairment Scale; Ditunno et al., 1994)

Result of Trauma	Treatment Measures
Neck sprain/whiplash in- jury	Analgesics, application of heat/cold, immobilization (as brief as possible). Early in- itiation of active exercise therapy. Measures to prevent chronification
Fracture	Stable \Rightarrow conservative (extension/fixation). Unstable \Rightarrow surgery
Arterial dissection	Anticoagulation
Spinal cord trauma	Methylprednisolone (i. v.) within 8 hours of trauma (bolus of 30 mg/kg over 15 min, then 5.4 mg/kg/h for 23 hours). Monitor respiratory and cardiovascular function, bladder/bowel function; thrombosis prophylaxis, pain therapy, careful patient positioning and pressure sore prevention. Transfer to specialized center for rehabilitation of paraplegic patients (as indicated)

Table 38 Treatment of spinal trauma (p. 274)

Features	Site of Lesion	Clinical Manifestations
Spinal cord transection (p. 48)	 Cervical spinal cord Thoracic spinal cord Lumbar/sacral spinal cord 	 Quadriplegia Paraplegia Paraplegia/conus syndrome with paralysis of blad- der/rectum and saddle anesthesia
Lesion affecting a por- tion of the spinal cord (pp. 32, 50)	 Anterior root Posterior root Incomplete transverse cord syndrome Complete transverse cord syndrome 	 Flaccid paralysis, muscle atrophy, hyporeflexia (⇒ segment-indicating muscles, see Table 2, p. 357) Localized/radicular/referred pain, sensory deficit in corresponding dermatome Brown-Séquard syndrome, posterior column syndrome, anterior horn syndrome, posterior horn syndrome, central cord syndrome, anterior spinal syndrome See p. 48
Temporal course	AcuteChronic	Spinal shockSpasticity, sensory and autonomic dysfunction

 Table 39
 Clinical manifestations of spinal cord lesions (p. 282)

 Table 40
 Malformations and developmental anomalies (p. 288)

Feature	Syndrome ¹	Comments ²
Macrocephaly (ab- normally large head)	Hydrocephalus (p. 290), hydranencephaly, megalencephaly (massively enlarged brain)	4th week/ 2nd to 4th month
Craniostenosis (premature ossifi- cation of cranial sutures, p. 4)	Turricephaly (\Rightarrow lambdoid and coronal suture; oxycephaly), scaphocephaly (\Rightarrow sagittal suture; dolichocephaly, "long head"), brachycephaly (\Rightarrow coronal suture; "short head")	Before 4th year of life
Migration disorder (defective migra- tion of neuroblasts into cortex)	Schizencephaly (presence of cysts or cavities in the brain), agyria (lissencephaly ² , few or no convolutions), pachygyria (broad, plump convolutions), heterotopia/dystopia (ectopic gray matter)	2nd to 5th month
Microcephaly (ab- normally small head)	Micrencephaly (abnormally small brain)	5th week (primary), peri- or postnatal (sec- ondary)
Dysraphism (neural tube defect)	See p. 292	3rd to 4th week/ 4th to 7th week
Chromosomal anomaly	Down syndrome (trisomy 21, mongolism), Patau syndrome (trisomy 13), Edwards syndrome (trisomy 18), cri-du-chat syndrome (deletion, short arm of chromosome 5), Klinefelter syndrome (XXY), Turner syndrome (XO), fragile-X syndrome	Genome mutation
Phakomatosis	See p. 294	
Prenatal or peri- natal infection	Rubella, cytomegalovirus, congenital neurosyphilis, HIV/ AIDS, toxoplasmosis	
Mental retardation	A component of many syndromes (e.g., microcephaly, hy- drocephalus, Down syndrome, perinatal or prenatal infec- tion)	
Cerebral lesion	Ulegyria (postanoxic corticomedullary scarring), poren- cephaly (p. 290), hemiatrophy, infantile cerebral palsy (p. 288 f)	Prenatal, perinatal or postnatal

1 (Selected). **2** The times specified refer to the gestational and neonatal ages, respectively. **3** There are two forms of lissencephaly: type 1, Miller–Dicker syndrome (craniofacial deformity), and type 2 (pronounced heterotopia with Fukuyama muscular dystrophy).

381

Table 41 Age-related changes (p. 296)

Change	Sequelae	Elevated risk ¹ of
Accommodation	Presbyopia	
Miosis	I Light/convergence reaction	
Cataract	Glare, 🌐 visual acuity	Blindness
I Hearing (inner ear)	Presbycusis	Deafness
Sense of smell/taste	Impaired sense of smell/taste	
û Body fat	$\hat{\mathbb{T}}$ volume of distribution for fat-soluble drugs²	Obesity
 Total body water, Thirst 	$\ensuremath{\mathbbm I}$ volume of distribution for water-soluble drugs^2	Dehydration, hydro- penia
Arteries	Atherosclerosis, impairment of cerebral autoregula- tion and blood-brain barrier, decrease in cerebral blood flow, reduced tolerance of brain tissue to ischemia and metabolic changes	Stroke ³ , leukoaraiosis ⁴ , subcortical arterios- clerotic encephalo- pathy (p. 172), cerebral amyloid angiopathy ⁵ , atrial fibrillation, myo- cardial infarction
Motor function	Dobility, Treactivity, coordination, fine motor control, muscle atrophy (especially thenar, dorsal in- terosseous, and anterior tibial muscles), U muscle force, file g muscle tone, hypokinesis of arms, gait impairment (p. 60)	Falls (p. 204), osteo- porosis, fear of falling/ inactivity (avoidance of social contact, isola- tion)
Reflexes	Reflex movements (p. 42), palmomental reflex, snout reflex, grasp reflex	Falls
Sensation	Pallhypesthesia in toe/knuckle region, ${\mathbb Q}$ position sense	Polyneuropathy, ataxia, falls
Brain atrophy	Senile forgetfulness ⁶ (impairment of episodic memory, p. 134)	Alzheimer disease, leukoaraiosis ⁴
$\ensuremath{\mathbb{Q}}$ Cerebral dopamine synthesis	Stooped posture	Parkinsonism
Cerebral norepinephrine		Depression
D Non-REM stage 4 (p. 112)	Early awakening, insomnia	Sleep apnea syndrome

(Resnick, 1998)

1 Risk of developing condition in old age. 2 Increased risk of drug side effects. 3 Especially due to border zone infarction, subdural hematoma (after relatively minor trauma). 4 Rarefaction of white matter seen as bilateral, usually symmetrical hypodensity on CT and as hyperintensity on T2-weighted MRI (FLAIR = fluid-attenuated inversion recovery sequence). 5 Increased risk of spontaneous intracranial hemorrhage (p. 176). 6 Benign senescent forgetfulness, age-associated memory impairment (AAMI).

 Table 42
 Criteria for differentiation between dementia and depression (p. 297)

Dementia	Depression
Patients seem indifferent to memory impairment; semantic paraphasia	Patients describe memory impairment precisely and in detail
Tests reveal cognitive deficit	Tests reveal minimal or no cognitive deficit
Depressive manifestations develop slowly (second- ary)	Depressive manifestations prominent on presenta- tion (brooding, anxiety, early awakening, loss of ap- petite, self-doubt)
Rare past history of depression	Frequent past history of depression

 Table 43
 Criteria for differentiation between different types of hyperkinesia (p. 300)

Syndrome	Features
Chorea (p. 66)	Overshooting, spontaneous, abrupt, alternating, irregular movements. Promi- nence varies from restlessness with little gesticulation, fidgety hand movements and hesitant, dance-like gait impairment to continuous, flowing, violent, disabling hyperkinesias
Dystonia (p. 64)	Involuntary, continuous and stereotyped muscle contractions that lead to rotat- ing movements and abnormal posture
Athetosis (p. 66)	Localized peripheral dystonic movements
Ballismus (p. 66)	Violent, mainly proximal flinging movements of the limbs
Tics (p. 68)	Repetitive, stereotyped, localized twitches that can be voluntarily suppressed, but with a build-up of inner tension
Myoclonus (p. 68)	Brief, sudden, shocklike muscle twitches occurring repetitively in the same muscle group(s)

(Harper, 1996)

 Table 44
 Symptomatic forms of parkinsonism (p. 302)

Cause	Examples
Infectious disease	Encephalitis lethargica' (von Economo postencephalitic parkinsonism), measles, tick-borne encephalitis, poliomyelitis, cytomegalovirus, influenza A, herpes sim- plex
Intoxication	MPTP2, manganese (miners, industrial workers), carbon monoxide, methanol
Drugs ³	Neuroleptics (phenothiazines ⁴ , butyrophenones ⁵ , thioxanthenes ⁶ , benzamide ⁷), reserpine, calcium channel blockers (cinnarizine, flunarizine)
Other diseases ⁸	Multiple brain infarcts/subcortical arteriosclerotic encephalopathy ⁹ , punch-drunk encephalopathy (dementia pugilistica), normal pressure hydrocephalus (p. 160), brain tumor (frontal), subdural hematoma, calcification of basal ganglia ¹⁰ , neu- roleptic malignant syndrome ¹¹

1 In the aftermath of the influenza pandemic that followed the First World War (now only of historical interest). 2 1-Methyl-4-phenyl-1,2,3,6-tetrahydropyridine. MPTP is converted into MPP⁺, which accumulates in dopaminergic neurons and interferes with electron transfer in the mitochondrial respiratory chain, leading to an accumulation of free radicals and neuronal death. An outbreak of MPTP-induced parkinsonism occurred in California in the early 1980s, when this substance appeared as a contaminant of opiate drugs synthesized in clandestine laboratories for illegal use. 3 Parkinsonoid. 4 Fluphenazine, levomepromazine, peraine, perphenazine, promazine, triflupromazine, etc. 5 Benperidol, fluspirilene (diphenylbutylpiperidine), haloperidol, etc. 6 Chlorprothixene, clopenthixol, fluanxol. 7 Metoclopranide. 8 Includes the terms pseudoparkinsonism, hypokinetic-rigid syndrome, and hypertonic-hyperkinetic syndrome. 9 Lower-body parkinsonism. 10 Autosomal recessive (Fahr disease), associated with hypoparathyroidism and pseudohypoparathyroidism. 11 Parkinsonian hyperthermia syndrome: rigidity, hyperthermia, impairment of consciousness; induced by neuroleptic drugs or by the use or withdrawal of levodopa or other dopaminergic agonists.

Syndrome	Features		
Hereditary			
Familial spastic paraple- gia (SPG, p. 286)	Uncomplicated SPG1-2: SPG26 (X/Xq22/PLP = proteolipid protein), SPG3A (AD/ 14q11.2-24.3/atlastin), SPG46 (AD/2p22-p21/spastin), SPG5A (AR/8p12-q13/?), SPG6 (AD/15q11.1/?), SPG76 (AR/16q24.3/paraplegin), SPG8/AD/8q23-24/?), SPC10 (AD/12q13/?), SPG11 (AR/15q13-15/?), SPG12 (AD/19q13/?), SPG13 (AD/ 2q24-34/?) Complicated SPG3 ⁻ SPG1 ⁶ (X/Xq28/L1-CAM = L1 cell adhesion molecule), SPG2 ⁶ (X/Xq22/PLP), SPG7 ⁶ (AR/16q24.3/paraplegin), SPG9 (AD/10q23.3-24.1/?), SPG14 (AR/3q27-28/?), SPG16 (X/Xq11.2/?)		
Adrenomyelo- neuropathy ⁴	X-linked recessive, onset usually after age 20. Progressive spastic paraparesis, polyneuropathy, urinary incontinence, sometimes hypocortisolism. A similar syndrome develops in 20% of all female heterozygotes (carriers)		
Spinocerebellar ataxia type 3	See p. 280		
Acquired			
Primary lateral sclerosis	Onset usually after age 50. Slowly progressive symmetrical paraspasticity without marked weakness, dysarthria. More common in men than in women		
Lathyrism (p. 304)	Onset usually before age 50. Subacute or chronic development of gait distur- bances (tip-toe/scissors gait, dorsal tilting of trunk), leg cramps, paresthesiae, uri- nary retention		
Tropical spastic para- paresis (TSP) ⁵	Onset: Slow up to age 60. Back pain, dysesthesiae, spastic paraparesis, urinary re- tention, impotence		

 Table 45
 Diseases affecting the first (upper) motor neuron (p. 304)

1 Abbreviations: AD = autosomal dominant, AR = autosomal recessive, X = X-chromosome/gene locus/gene product (? = unknown). **2** Isolated progressive spastic paraparesis. **3** As in uncomplicated SPG with additional manifestations including cerebellar ataxia, dystonia, optic neuropathy, tapetoretinal degeneration, muscle atrophy, dysarthria, deafness, sensory neuropathy, ichthyosis, dementia. **4** Impaired β -oxidation of very long chain fatty acids (VLCFA; C₂₄₋₂₆) due to defective peroxisomal transport \Rightarrow accumulation of VLFCA in nervous system, adrenal cortex, plasma. **5** HTLV-1-associated myelopathy (= HAM, human T-cell/lymphotropic virus type I); Jamaica, Japan, Caribbean. **6** Molecular genetic tests are available.

Table 46 Dis	seases affecting	the second	(lower)	motor neuron	(p. 304)
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Syndrome ¹	Features	
Hereditary		
Proximal ² (SMA) ³	SMA I ⁴ : onset < 3 months. AR. Flaccid quadriparesis ⁵ . Triangular mouth shape, paradoxical respiratory movements, impaired sucking ability, unable to sit SMA II: onset < 5 years. AR. The children learn to sit independently, but never to stand/walk. Scoliosis, joint contractures SMA III ⁶ : SMA IIIa: onset 3 years. AR. Delayed motor development. Children learn to stand/walk SMA IIIb: onset 3-30 years. AR. Development normal. Calf (pseudo)hypertrophy. Absence of bulbar muscle involvement. CK ⁷ sometimes elevated SMA IV: onset > 30 years. AR	
Nonproximal [®] SMA	Distal SMA ⁹ : forms with different ages of onset (infantile, juvenile, adult). Usually slow course, sometimes stabilizing after a few years, sometimes progressive. May be accompanied by myoclonus, deafness, dysphonia, dysarthria, and/or ataxia. Scapuloperoneal muscular atrophy ¹⁰ : Onset: Adolescence or adulthood. Weakness ⇔ foot dorsiflexors, shoulder girdle, arm Progressive bulbar palsy ¹¹ : Onset in adulthood ¹² . Progressive weakness of bulbar muscles. Muscular atrophy and respiratory muscle involvement develop as the disease progresses	
Spinobulbar muscular atrophy (Kennedy type)	Onset: 20–70 years. Gynecomastia, gradual progression of muscular atrophy (legs > arms, proximal > distal, asymmetrical; dysarthria, dysphagia, tongue atrophy). Slight elevation of CK. Gene locus Xq12	
Acquired		
Acute viral infection	Poliomyelitis (p. 242), other enteroviruses (e. g., echovirus, coxsackievirus, enter- ovirus type 70/71 \Rightarrow acute hemorrhagic conjunctivitis), mumps virus	
Postpolio syndrome	See p. 242	
Lymphoma	Accompanies Hodgkin and non-Hodgkin lymphomas. Elevation of CSF protein, oligoclonal IgG in CSF	
Radiation-induced	Develops months to years after irradiation of para-aortic lymph nodes (testicular tumors, uterine carcinoma). May progress rapidly	

(Tandan, 1996; Rudnik-Schöneborn, Mortier and Zerres, 1998)

AR = autosomal recessive, AD = autosomal dominant; XR = X-linked recessive. 1 Selected syndromes. 2 Proximal muscle involvement. 3 SMA = spinal muscular atrophy; gene locus for SMA I to III Sq12.-q13.3 4 AR infantile SMA, Werdnig-Hoffmann disease. 5 Floppy baby syndrome, froglike posture in supine position; lack of head control. 6 AR > AD; juvenile SMA; Kugelberg-Welander disease. 7 Creatine kinase (CG). 8 Distal or localized (monomelic segmental SMA) muscle groups, symmetrical or asymmetrical involvement. 9 AP > AD. 10 AD \simeq Onset age 30-90 years, slowly progressive; AR \simeq onset < 5 years; may progress slowly. 11 AD/AR. 12 Fazio-Londe type with onset at age 2–13 years, rapidly progressive

Diagnostic Categories ¹			
Definite ALS ²	Evidence of first + second motor neuron lesion in 3 regions of the body ³		
Probable ALS	Evidence of first + second motor neuron lesion in 2 regions of the body		
Possible ALS	Evidence of first + second motor neuron lesion in 1 region of the body or evi- dence of first motor neuron lesion in 2 to 3 regions of the body		
Suspected ALS	Evidence of second motor neuron lesion in 2 to 3 regions of the body		
Diagnostic features	Progressive symptoms of a first (p. 46) and second motor neuron lesion (p. 50). Fasciculation in more than one region of the body. Neurogenic EMG findings, nor- mal nerve conduction velocity/absence of motor conduction block. Absence of sensory deficits, sphincter dysfunction, visual disturbances, autonomic dysfunc- tion, parkinsonism, and Alzheimer, Pick or Huntington disease		
Syndromes with manife- stations similar to those of ALS	Cervical radicular syndromes, cervical myelopathy, monoclonal gammopathy. Multifocal motor neuropathy (GM ₁ antibodies). Lymphoma; paraneoplastic syn- drome; hyperthyroidism, hyperparathyroidism; diabetic amyotrophy: postpolio syndrome; hexosamidinase A deficiency. Radiation-induced lesion. Toxicity (lead, mercury, manganese). Myopathy (inclusion body myositis, polymyositis, muscular dystrophy). Spinal muscular atrophy. Creutzfeldt–Jakob disease		

 Table 47
 Diseases affecting both the first (upper) and the second (lower) motor neurons (p. 304)

1 According to Leigh and Ray-Chaudhuri (1994). 2 ALS = amyotrophic lateral sclerosis. 3 Brain stem, proximal/distal arm, chest, proximal/distal leg.

Syndrome	Defect/Enzyme Defect	Symptoms and Signs
Galactosemia	Galactose-1-phosphate uridyltransferase ¹	Milk intolerance, apathy, jaundice, anemia, cata- ract, psychomotor retardation
Nonketonic hyperglycine- mia	Defective conversion of glycine to serine	Hypotonia, dyspnea, myoclonus, generalized seizures
Hyperammonemia	Urea cycle ²	Crisislike episodes of vomiting, sucking weakness, somnolence, coma, seizures, hyperpnea, hyper- pyrexia
Maple syrup urine disease	Defective breakdown of branched-chain amino acids	Hypotonia, seizures, coma, ketoacidosis
Zellweger syndrome	Peroxisomes ³	Hypotonia, sucking weakness, nystagmus, seizures, craniofacial dysmorphism

1 Multiple types — high galactose-1-phosphate levels. 2 Defects of all six enzymes of the urea cycle are known. Adult onset is rare. Hyperammonemia in defects of carbamoyl-phosphate synthetase, ornithine carbamoyltransferase, argininosuccinic acid synthetase (citrullinemia), argininosuccinase. Arginase defect leads to arginemia. 3 Cytoplasmic organelles that mediate fatty acid oxidation, bilary acid and cholesterol synthesis, pipecolic and phytanic acid metabolism, and plasmalogen (myelin) synthesis. Other peroxisomal syndromes: neonatal adrenoleukodystrophy, infantile Refsum disease, hyperpipecolatemia.

Syndrome	Defect/Enzyme Defect	Symptoms and Signs
Tay–Sachs disease ¹	Hexosaminidase A (⇔ ac- cumulation of ganglioside GM₂)	Abnormal acoustic startle reaction, delayed development. Begins with muscular hypotonia, followed by spasticity, seizures, blindness, dementia, and optic nerve atrophy ²
Gaucher disease ³ (type II, acute neuropathic)	Glucocerebrosidase (⇔ lipid storage)	Loss of motor control, apathy, dysphagia, retro- flexion of the head, strabismus, splenomegaly
Niemann–Pick disease ⁴ (type A)	Sphingomyelinase defi- ciency (sphingomyelin storage ⇔ Niemann–Pick cells)	Enlargement of spleen, liver and lymph nodes; pulmonary infiltrates, spasticity, muscular axial hypotonia, blindness, nystagmus, macular cherry red spot
GM ₁ gangliosidosis	β-Galactosidase	Craniofacial dysmorphism. Initial flaccid paralysis that later becomes spastic; loss of visual acuity, nystagmus, strabismus, hepatomegaly
Krabbe disease (globoid cell leukodystrophy)	β-Galactocerebrosidase (galactocerebroside û)	General muscular hypertonia, vomiting, opistho- tonus, spasticity, blindness, deafness
Pelizaeus-Merzbacher disease ⁵	Proteolipid protein synthe- sis	Nystagmus, ataxia, psychomotor retardation, choreoathetosis

 Table 49
 Metabolic encephalopathies of infancy (first year of life, p. 306)

1 GM₂ gangliosidosis. **2** Cherry red spot in optic fundus is found in over 90% of cases. **3** Glucocerebrosidase; three known subtypes \Rightarrow type 1: nonneuropathic (juvenile) form with hematological changes and bone fractures; type 3: see p. 307. **4** Different types (A, B, C) exist. Type B does not produce neurological symptoms. **5** Other sudanophil (orthochromatic) forms of leukodystrophy are known.

Table 50 Stages of hepatic/portosystemic encephalopathy (p. 308)

Stage	Behavior ¹	Motor function	EEG ²
1 I	Attention deficit, impaired concentration, euphoria or de- pression, dysarthria, insomnia	Handwriting illegible, asterixis +/-	Usually normal to $\boldsymbol{\theta}$ waves
Ш	Sleepy, marked behavioral changes (confusion, disorien- tation, apraxia)	Asterixis	Pathological (δ waves)
Ш	Severely confused, somnolent to soporific	Asterixis	Pathological (δ/triphasic waves)
IV	Coma	Presence (stage IVa) or ab- sence (stage IVb) of motor re- sponses to pain	Pathological (triphasic/ arrhythmic ð/sub-ð waves)

(Adams and Foley, 1952)

1 Earlier stages are assessed with psychometric methods, e.g., number connection test (time required for patient to connect 25 numbered circles in numerical order), ability to draw a five-pointed star. 2 Nonspecific changes; actual findings may differ.

Appendix

Site	Syndrome ⇔ Time course	Symptoms and Signs	Common Tumors	Lesions/Antibodies
Cerebrum	Photoreceptor/retinal degeneration ⇒ weeks to months	Progressive blindness without pain	Small-cell lung cancer	Loss of photorecep- tors/anti-CAR ¹
	Limbic encephalitis ⇒ weeks to months	Restlessness, confu- sion, memory impair- ment	Small-cell lung cancer	Neuronal loss, peri- vascular and mening- eal lymphocytic infil- trates ⇒ medial tem- poral lobe, limbic sys- tem/ANNA-1 ²
Cerebel- lum, brain stem	Brain stem encephali- tis ⇔ days to weeks	Dysphagia, dys- arthria, nystagmus, diplopia, ataxia, dizzi- ness	Small-cell lung cancer	Neuronal loss, inflam- matory infiltrates in the brain stem
	Subacute cerebellar degeneration \Rightarrow weeks to months	Cerebellar ataxia, dysarthria, nystag- mus, diplopia, ver-	Small-cell lung cancer, carcinoma of ovary/breast, Hodg- kin disease	Death of Purkinje cells/APCA ³
	Opsoclonus-myo- clonus ⇔ weeks	tigo Abrupt, irregular eye and muscle move- ments, cerebellar ataxia, encephalo- pathy	Neuroblastoma (children); breast cancer/lung cancer (adults)	Neuronal loss in den- tate nucleus (adults)/ANNA-2 ⁴
Spinal cord	Necrotizing my- elopathy ⇔ hours, days to weeks	Flaccid para-/ quadri- paresis, bladder/ bowel dysfunction, segmental sensory loss	Small-cell lung cancer, lymphoma	Necrosis of white and gray matter of spinal cord
	Stiff man syndrome⁵ ⇔ days to weeks	Painful muscular rigidity initially trig- gered by emotional, acoustic and/or tac- tile stimuli; auton- omic dysfunction	Small-cell lung cancer, Hodgkin lym- phoma, breast cancer, pharyngeal carcinoma	Transitory high-cervi- cal lesions may be seen in MRI scans/ anti-GAD ⁶

Table 51	Paraneoplastic	syndromes	of the	CNS	(n 312)
Table 31	rarancopiastic	syndionics	or the	CIVD	(p. 512)

(Brown, 1998)

1 CAR = cancer-associated retinopathy. 2 ANNA-1 = antineural nuclear antibody type 1 = anti-Hu. 3 APCA = anti-Purkinje cell cytoplasmic antibody = anti-Yo. 4 ANNA-2 = antineural nuclear antibody type 2 = anti-Ri. 5 Manifestations variable (e. g., axial or distal muscle may be more prominent); attributed to diminished supraspinal inhibition of motor neurons, leading to continuous contraction of agonists and antagonists. 6 Anti-GAD = glutamic acid decarboxylase antibodies; antibodies directed against amphiphysin (terminal synaptic protein) have also been found.

Substance	Adverse Effects1
Neuroleptics	Drug-induced parkinsonism (p. 383, Table 44), early/late dyskinesia (p. 66), akathisia², low seizure threshold
Antidepressants	Somnolence, increased drive, confusion, akathisia, low seizure threshold, tremor, serotonin syndrome $^{\rm 3}$
Aspirin ⁴	Tinnitus, dizziness
Baclofen	Fatigue, depression, headaches, low seizure threshold
Levodopa, dopamine agonists	Confusion, hallucinations, psychosis, insomnia, hyperkinesia
Corticosteroids, ACTH Antibiotics	Depression, increased drive, mania, insomnia, headaches, dizziness, sweating, low seizure threshold, tremor <i>Aminoglycosides</i> : Tinnitus, hearing impairment. <i>Quinolone derivatives</i> : Insomnia, hallucinations, headaches, low seizure threshold, dizziness, somnolence, tinnitus. <i>Tetracyclines</i> : Pseudotumor cerebri (children), abducens paralysis (adults)
Glycosides	Visual disturbances, somnolence, hallucinations, seizures, delirium
Calcium antagonists	Fatigue, insomnia, headaches, depression. <i>Flunarizine/cinnarizine</i> : Drug-induced parkinsonism
Coumarins	Intracranial hemorrhage (2–12%/year)
Radiotherapy ⁵	Acute (<1 week): Headaches, nausea, somnolence, fever. Subacute: (2–16 weeks): Somnolence, focal neurological deficits, leukoencephalopathy, brain stem syn- drome (rare). Late (>4 months): radiation necrosis ⁶ , leukoencephalopathy, dementia, secondary tumor
Chemotherapy ⁷	Acute: Insomnia, confusion, restlessness, stupor, generalized seizures, myoclonus. Late: Apathy, dementia, insomnia, incontinence, gait impairment, ataxia

Table 52 latrogenic encephalopathies (p. 314)

(Biller, 1998; Diener and Kastrup, 1998; Keime-Guibert et al., 1998)

1 Common adverse effects. 2 Inability to sit still with tormenting sensations in the legs that improve briefly when the patient moves about. 3 Characterized by confusion, fever, restlessness, myoclonus, diaphoresis, tremor, diarrhea, and ataxia; usually due to drug interactions, e. g., fluoxetine + sertraline, serotonin reuptake inhibitor + tryptophan, MAO inhibitors, carbamazepine, lithium, or clomipramine. 4 Acetylsalicylic acid (ASA). 5 Syndromes also occur in combination with chemotherapy. 6 One to two years after percutaneous radiotherapy, ca. 6 months after interstitial radiotherapy. Focal neurological deficits. 7 Methotrexate (high-dose i. v., intrathecal) cisplatin, vincristine, asparaginase, procarbazine, 5-fluorouracil, cytosine arabinoside, nitrosourea compounds (high-dose), ifosfamide, tamoxifen, etoposide (high-dose).

Table 53 Neuropathy syndromes (p. 316)

Syndrome	Causes
Predominantly sym- metrical motor deficits	Amyotrophic lateral sclerosis (ALS), multifocal motor neuropathy, Guillain-Barré syndrome, CIDP ¹ , acute porphyria, hereditary sensorimotor neuropathy
Predominantly asym- metrical or focal motor deficits	Neuronopathy: ALS, poliomyelitis, spinal muscular atrophy Radicular lesion: Root compression (herniated intervertebral disk, tumor), herpes zoster, carcinomatous meningitis, diabetes mellitus Plexus lesion: Neuralgic amyotrophy of shoulder, tumor infiltration, diabetes melli- tus, tomaculous neuropathy, compression (positional) Multiple mononeuropathy: Vasculitis, diabetes mellitus, multifocal motor neu- ropathy, neuroborreliosis, sarcoidosis, HIV, tomaculous neuropathy, leprosy, neu- rofibromatosis, cryoglobulinemia, HNPP (p. 332), neoplastic infiltration Mononeuropathy: Compartment syndrome (median n., ulnar n.), compression (anterior interosseous n., peroneal n.), lead poisoning, diabetes mellitus
Predominantly auton- omic disturbances	Diabetes mellitus, amyloidosis, Guillain-Barré syndrome, vincristine, porphyria, HIV, idiopathic pandysautonomia, botulism, paraneoplastic neuropathy
Predominant pain	Diabetes mellitus, vasculitis, Guillain–Barré syndrome, uremia, amyloidosis, ar- senic, thallium, HIV, Fabry disease, cryptogenic neuropathy
Predominantly sensory disturbances	Diabetes mellitus, alcohol, ethambutol, vitamin B_{12} deficiency, folic acid deficiency, overdosage of vitamin B_{6} , paraneoplastic, metronidazole, phenytoin, thalidomide, leprosy, cytostatic agents (e.g., vincristine, vinblastine, vindesine, cisplatin, pacitiaxel), amyloidosis (dissociated sensory loss), hereditary sensory neuropathy, monoclonal gammopathy, tabes dorsalis, Friedreich ataxia (p. 280)
Ganglioneuropathy² (ataxia)	Paraneoplastic, Sjögren syndrome, cisplatin, vitamin ${\rm B}_6$ intoxication, HIV, idiopathic sensory neuronopathy

(Barohn, 1998)

1 Chronic inflammatory demyelinating polyneuropathy. 2 Asymmetrical proprioceptive loss without paralysis.

Table 34 Acquired and herearchy hearopathies (p. 510)					
Cause	Examples				
Acquired • Metabolic disorder • Dietary deficiency • Immune-mediated	 Diabetes mellitus, uremia, hypothyroidism, acromegaly Vitamin deficiency (B₁ = beriberi, B₆, B₁₂, E), malabsorption Guillain–Barré syndrome, Fisher syndrome, chronic inflammatory demyelinating polyneuropathy (CIDP), multifocal motor neuropathy, pandysautonomia, neuralgic amyotrophy of shoulder¹, vasculitis, connective tissue disease, plasmocytoma, benign monoclonal gammopathy, Churg–Strauss syndrome, cryoglobulinemia, rheumatoid arthritis 				
Infection	 Herpes zoster, leprosy, Lyme disease, HIV, neurosyphilis, diphtheria, typhus, paratyphus 				
• Drugs	 Carbimazole, cisplatin, cytarabine, enalapril, ethambutol, etoposide, gentami- cin, gold, imipramine, indomethacin, INH, paclitaxel, phenytoin, procarbazine, suramine, thalidomide, vinca alkaloids, vitamin B₆ 				
 Toxins (environmen- tal, industrial)², drugs² Neoplasm Mechanical Unknown 	 Alcohol, arsenic, benzene, lead, heroin, hexachlorophene, pentachlorophenol, polychlorinated biphenyls, mercury, carbon tetrachloride, thallium, triarylophosphate Paraneoplastic: Lung, stomach, or breast cancer; Hodgkin disease, leukemia. Infiltration: Hodgkin disease, leukemia, carcinomatous meningitis, polycythemia Compression, trauma, distortion Critical illness polyneuropathy 				
Hereditary	See pp. 332 and 396 f				

Table 54 Acquired and hereditary neuropathies (p. 316)

390

(Barohn, 1998)

1 Causes not confirmed. 2 A large number of substances can lead to polyneuropathy (PNP). Only some of them are listed.

Table 55	Additional	diagnostic	studies for	r neuropathies	(p. 316)

Method	Information/Parameters
Neurography	Motor neuron lesion: Normal (consider conduction block) Ganglionopathy: MSAP ¹ normal to[], SNAP ² normal to[], (dermatomal) SEP ³ [] Radiculopathy: H reflex: Lateral inequality/absence ⁴ ; F waves: some prolonged; (dermatomal) SEP [] Axonal lesion: Motor NCV ⁵ : Normal; MSAP[], SNAP[] Demyelination: DML ^Q] NCV:[] or local conduction block (localize by inching); MSAP[] (dispersed; F waves: Prolongation or absence; SNAP: normal/dispersed in motor neuropathy, [] in sensory neuropathy
Needle electro- myography	Motor neuron lesion: Fibrillation, positive waves, fasciculation. Amplitude, poly- phasia rate, MAP ² duration ¹ Ganglionopathy: MAP ¹ Low-grade neurogenic changes may be observed <i>Radiculopathy</i> : Pathological spontaneous activity in paravertebral muscles/seg- ment-indicating muscles (p. 357); MAP ⁸ : neurogenic changes <i>Axonal</i> lesion: Pathological spontaneous activity (fibrillation, fasciculation); MAP: neurogenic changes <i>Demyelination</i> : Absence of pathological spontaneous activity, maximum innerva- tion with thinning pattern
Laboratory tests ⁹	Standard tests: Erythrocyte sedimentation rate, differential blood count, blood glucose (diurnal profile), C-reactive protein, calcium, sodium potassium, alkaline phosphatase, SGOT ¹⁰ , SGPT ¹¹ , CK ¹² , γ -GT ¹³ , electrophoresis, rheumatoid factors, vitamin B ₁ / folic acid, <i>Borrelia</i> /HIV antibodies, basal TSH ¹⁴ , triglycerides, cholesterol, urine status, blood culture <i>Special tests</i> : CSF, homocysteine, hemoglobin A _{1C} , syphilis serology, parathyroid hormone, antinuclear antibodies (e.g., Sm, RNP, Ro SS-A, La SS-B, Scl-70, Jo-1, Pm-Scl), antineuronal antibodies (ANNA-1, anti-Hu), myelin-associated glycoprotein (MAG), ganglioside antibodies (GM ₁ , GD _{1a} , GD _{1b} , GQ _{1b}), heavy metals (blood, urine), porphyrins, cryoglobulins, serum phytanic acid, very long chain fatty acids (VLCFA, C ₂₄₋₂₆), molecular genetic testing
Sural nerve biopsy ¹⁵	Vasculitis, amyloid neuropathy, neuropathy with sarcoidosis, leprosy, chronic neuropathy (HMSN III/metachromatic leukodystrophy, with or without other forms of hereditary neuropathy \Rightarrow p. 332; chronic inflammatory neuropathy, polyglucosan body neuropathy), tumor (neurofibroma/schwannoma, neoplastic infiltration; paraneoplastic neuropathy), neuropathy with monoclonal gammopathy, if applicable
Diagnostic imaging	Guided by clinical findings (spinal, radicular, plexus, distal peripheral lesions?), plain radiographs, ultrasound, CT, MRI, myelography, skeletal scintigraphy and/or angiography

 1 = elevated, prolonged;
 0 = diminished, absent
 1 Summated muscle action potential (evoked).
 2 Amplitude of sensory nerve action potential.
 3 Somatosensory evoked action potential amplitude reduced or absent.
 4 For technical reasons, can only be determined for \$1. 5 Nerve conduction velocity. 6 Distal motor latency. 7 Muscle action potential. 8 After 2 weeks, at earliest. 9 Par-tial list. 10 Serum glutamic–oxaloacetic transaminase. 11 Serum glutamate–pyruvate transaminase. 12 Creatine kinase. 13 γ-Glutamyl transpepsidase. 14 Thyrotropin. 15 Muscle biopsy may also be helpful in some cases.

Table 56	Causes	of	radicular	S	/ndromes	(p.	. 318))
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Cause	Comments
 Degenerative changes Intervertebral disk herniation Spondylosis deformans Spinal canal stenosis Spondylolisthesis 	 Symptoms usually resolve with conservative treatment¹ Torus-, buckle- or spur-shaped spondylophytes form due to degenerative changes in the intervertebral disks See p. 284 Slippage of a vertebra with respect to the next lower vertebra because of bilateral spondylolysis²
Trauma	See p. 272
Neoplasm	Primary spinal tumor (p. 284), metastatic tumor/neoplastic meningeosis (p. 262)
Infection/inflammation	See p. 222 ff. herpes zoster, borreliosis, epidural abscess, spondylitis, sar- coidosis, arachnopathy
Vascular	See p. 282
Metabolic	Diabetes mellitus (p. 324; Table 59, p. 395)
Inflammatory rheumatic	Ankylosing spondylitis, rheumatoid arthritis
Malformation	See p. 288 ff
latrogenic	Injection, lumbar puncture, surgery
Radiotherapy	Radiation-induced amyotrophy (cauda equina) ³
Pseudoradicular syndrome4, nonradicular pain	 Arm: Carpal tunnel syndrome, stiff shoulder, humeroscapular periar- thropathy, syringomyelia Leg: Facet syndrome, sacroiliac joint syndrome, coccygodynia, coxarthro- sis, heterotopic ossification General: Polyradiculitis, connective tissue diseases, rheumatoid diseases, malformations, myopathy/muscle trauma, strain, arthropathy, en- dometriosis, osteomyelitis, osteoporosis, arterial dissection, prostatitis, cystitis, Paget disease, somatization disorder

(Mumenthaler et al., 1998)

1 Absolute indications for surgery: Massive lumbar disk herniation with sphincter dysfunction, cervical disk herniation with spinal cord compression, severe weakness. Relative indications: Persistent radicular pain, frequent recurrence of radicular symptoms. 2 Defect in the pars interarticularis of the vertebral arch. 3 Development of following manifestations months to years after radiotherapy (para-aortic irradiation): malignant testicular tumor, lymphoma; progressive flaccid paraparesis without major sensory loss. **4** To be considered in the differential diagnosis.

Table 57 Causes of plexus lesions (p. 318)

Cause	Comments
Neoplasm	 Upper limb: neurofibroma/schwannoma, metastatic tumor, breast/lung cancer (Pancoast tumor) Lower limb: urogenital tumors, cancer of the rectum, lymphoma
Vascular	Lower limb: psoas hematoma due to anticoagulation, hemophilia, aneurysm
Metabolic	Lower limb: diabetes mellitus (p. 395, Table 59)
Inflammatory	 Upper limb: neuralgic shoulder amyotrophy (p. 328) Lower limb: neuropathy of lumbosacral plexus, vasculitis
Trauma	 Upper limb: stab or gunshot wound, strain, contusion (trauma, birth), cervical nerve root avulsion (p. 272) Lower limb: pelvic fracture, sacral fracture
Compression	Upper limb: carrying heavy loads (backpack paralysis), thoracic outlet syndrome (T1–C8) ¹ , costoclavicular syndrome, hyperadduction syndrome
Infection	Lower limb: psoas abscess
latrogenic	 Upper limb: positioning, retraction (heart surgery), plexus anesthesia Lower limb: hip surgery, vascular surgery, hysterectomy, adverse positioning
Pregnancy	Lower limb: end of pregnancy, delivery
Radiotherapy	 Upper limb: brachial plexus paralysis months to years after radiotherapy Lower limb: due to radiotherapy of neoplasms in pelvic region

1 Thoracic outlet compression may be caused by a cervical rib, fibrous band or narrow scalene gap.

Table 58 Commo	on sites of mononeuropathy (pp. 318 and 312 †) Lesion ⇒ Syndrome	Cause ¹
		Dislocation of shoulder
Axillary nerve	Abduction paralysis, deltoid atrophy	
Long thoracic nerve	Winging of scapula; no sensory deficit, weakness of arm elevation	Compression ("backpack" paraly- sis), neuralgic shoulder amyo- trophy, postinfectious
Radial nerve	 UA² ⇒ hand drop with sensory ↓ in radial back of hand; prominent between 1st and 2nd finger PFA⁴ ⇒ supinator syndrome⁵ 	 Compression³/fracture of shaft of humerus Fractured head of radius
Median nerve	 UA ⇒ monkey hand⁶ PFA ⇒ pronator teres syndrome, anterior interosseous syndrome⁸ 	Compression, fractureStrain, compression, fracture
	 DLA⁹ □ carpal tunnel syndrome, brachialgia, noc- turnal paresthesia¹⁰ 	 Compression, arteriovenous fistula/uremia, rheumatoid arthritis, pregnancy, diabetes mellitus, hypothyroidism, monoclonal gammopathy
Ulnar nerve	 UA ⇔ clawhand PFA ⇒ clawhand DLA ⇔ different types of paralysis 	 Supracondylar process Trauma, compression, arthrosis, habitual Compression, habitual
Lateral femoral cutaneous nerve	Meralgia paresthetica ¹²	Compression
Femoral nerve	 Proximal lesion ⇒ paralysis of knee extensors Intrapelvic lesion ⇒ additional paralysis of hip flexors (gait disturbance) 	 Psoas hematoma/abscess Surgery (hip surgery, hysterectomy), trauma
Sciatic nerve	Peroneal + tibial (partial) lesion	 Trauma, hip surgery, intra- gluteal injection
Common per- oneal nerve	 Lesion at head of fibula ⇒ paralysis of dorsiflexors of foot (step gait) 	Compression, fracture, sprain, compartment syndrome
Tibial nerve	 Popliteal lesion ⇒ paralysis of all flexor muscles (foot, toes), sensory loss on back of calf and sole of foot; pain, absence of ankle jerk reflex 	• Fracture, compression
	 Lesion of lower leg ⇒ clawed toes; preservation of reflex Tarsal tunnel syndrome¹³, pain. 	CompressionCompression, trauma
	reflex	

Table 58	Common sites	of mononeuropathy	/ (pp. 318 and 312 f)
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1 Selection. 2 Lesion at level of upper arm. 3 "Park bench paralysis", tourniquet (ischemia). 4 Lesion at level of elbow, proximal forearm. 5 Deep branch lesion: Pain on extensor surface of forearm, no sensory deficit, paralysis of long finger/thumb extensors with preservation of (radial) lifting of hand. 6 Paralysis of radial hand/finger flexors, pronators, thenar atrophy; sensory U in first 31/2 fingers, trophic disturbances. 7 Pain on outer surface of forearm. 8 Lesion: anterior interoseous n.; no sensory deficit; flexor weakness in distal segments of thumb, index and middle finger. 9 Lesion of distal forearm, wrist. 10 Painful nocturnal/morning paresthesia (in arm) with sensory loss, thenar atrophy, and paralysis as the condition progresses. 11 Overextension of finger at metacarpophalangeal joint, flexion of middle and distal phalanges, sensory deficit in ulnar 11/2 fingers. 12 Paresthesiae, pain in outer surface of thigh. 13 Lesion behind medial malleolus.

Table 59	Diabetic	polyneuropathy	syndromes	(p. 324)
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Syndrome	Features
 Symmetric distribution Diabetic polyneuropathy (DPN)¹ Diabetic pandysautonomia Small-fiber polyneuropathy (PNP) with weight loss Hypoglycemic PNP 	 Distal, primarily sensory, with or without pain. NCV² [] and/or respiration-dependent change in heart rate (p. 370) []. Areflexia³, pall-hypesthesia or pallanesthesia in toes, either initially or over time. Progression: increased sensory loss, impairment of position sense (sensory ataxia), paresthesiae, trophic changes, and paralysis. Autonomic dysfunction Cardiovascular⁴, gastrointestinal⁵, urogenital⁶, and skin⁷ changes. Usually associated with DPN, but isolated occurrence is possible Painful (burning, dull dragging pain, more prominent at night). Autonomic dysfunction. Relatively mild impairment of somatic sensation, vibration and position sense, muscle strength and reflexes Seen in insulinoma. May also be caused by recurrent hypoglycemia
Asymmetric distribution • Lumbosacral radicular neu- ropathy/plexus neuropathy ^{8,9} • Thoracolumbar radiculoneuropathy ⁹ • Compression syndromes • Cranial mononeuropathy ^{9,10}	 Rarely symmetrical. Intense pain radiating from low back to upper thigh. Weakness and atrophy of muscles innervated by the femoral nerve. Loss of quadriceps reflex. Minimal sensory loss Segmental beltlike pain distribution, sensory deficit, abdominal wall paralysis Carpal tunnel syndrome (Table 58), ulnar lesion at level of elbow III ⇒ acute, painful¹¹ ophthalmoplegia, usually without pupillary involvement. VII ⇒ acute, often painful paralysis of the peripheral type
	(Taular and Duals 1000)

(Taylor and Dyck, 1999)

1 Cannot be clinically distinguished from uremic neuropathy. 2 Nerve conduction velocity (sensory/motor).
3 Mainly the gastrocnemius reflex. 4 Resting tachycardia, fixed heart rate. 5 Gastroparesis, diarrhea, constipation, bilary stasis, fecal incontinence. 6 Urinary retention, erectile dysfunction, retrograde ejaculation. 7 Hypohidrosis, hyperkeratosis. 8 Also referred to as proximal diabetic neuropathy, diabetic amyotrophy, and femoral neuropathy.
9 Indicative of favorable prognosis (partial or complete remission). 10 Local infection (rhinocerebral mucormycosis, p. 246, otitis externa circumscripta) can cause cranial nerve deficits in patients with diabetes mellitus. 11 Periorbital, frontotemporal, hemispheric.

Table 60	Clinical spectrum	of Guillain-Barré	syndrome (p. 326)
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Syndrome	Features ¹
Acute inflammatory demyelinating poly- radiculoneuropathy (AIDP) ²	Perivenous lymphocytic infiltrates and demyelination. IgM/IgG antibodies ³ against GM ₁
Acute motor-sensory axonal neuropathy (AMSAN) ⁴	Pronounced paralysis, early muscular atrophy. Axonal degeneration. IgG antibodies directed against GM_1
Acute motor-axonal neuropathy (AMAN) ⁴	Pure motor neuropathy with axonal degeneration. IgG antibodies directed against GM_1 , GD_{1a} , GD_{1b}
Miller Fisher syndrome (MFS)	Diplopia (usually external ophthalmoplegia), ataxia, areflexia. IgG antibodies to $GQ_{1b}{}^5$

(Hahn, 1998)

The most common manifestations are listed. Other motor, sensory, or autonomic disturbances may also occur.
 The most common form in Europe, North America, and Australia. 3 Ganglioside antibodies. 4 Commonly occur in North China, Japan, and Mexico; rarely in Western countries. 5 Detected in over 95% of all patients; correlated with the course of the disease.

 Progressive paralysis in more than one limb Hyporeflexia or areflexia Progression lasting days to 4 weeks Symmetry (relative) of involvement Mild sensory distur- bances Cranial nerve involve- ment, especially VII Resolution 2-4 weeks after end of progres- sion Autonomic dysfunc- tion Initial absence of fever C5F protein 1² Typical neurophysio- logical findings³ Markedly asymmetri- cal involvement Initial or persistent bladder/bowel dys- function Granulocytes in CSF; cell count > 50 mononuclear cells/ mm³ Sharply localized sensory loss in the trunk region 	Necessary ¹	Supportive ¹	Doubtful ¹	Exclusion ¹
	in more than one limb • Hyporeflexia or	 days to 4 weeks Symmetry (relative) of involvement Mild sensory distur- bances Cranial nerve involve- ment, especially VII Resolution 2–4 weeks after end of progression Autonomic dysfunc- tion Initial absence of fever CSF protein Ω² Typical neurophysio- 	cal involvement Initial or persistent bladder/bowel dys- function Granulocytes in CSF; cell count > 50 mononuclear cells/ mm ³ Sharply localized sensory loss in the	asthenia, botulism, poliomyelitis, toxic neuropathy Porphyria Recent diphtheria Isolated sensory dis- turbances without

Table 61 Diagnostic criteria: Guillain-Barré syndrome (GBS) (p. 326)

(Asbury and Cornblath, 1990)

1 "Necessary" = prerequisite for diagnosis of GBS; "supportive" = supports the diagnosis; "doubtful" = GBS unlikely; "exclusion" = excludes the diagnosis of GBS. **2** May be normal initially, then rise in the course of the disease to several g/l; blood-brain barrier dysfunction; cell count < 10 mononuclear cells/mm³. **3** *Early phase*: Partial conduction block with reduced amplitude of evoked motor response potentials (proximal stimulation), loss of reflex and F-wave responses due to a proximal lesion; EMG recordings show reduced number of activatable motor units. *Later stages*: Variably reduced motor response potential; EMG shows mild to marked denervation (the less marked, the better the prognosis).

 Table 62
 Genetic features of hereditary polyneuropathy (p. 332)

Syndrome	Mode of Inheritance ¹ / Gene Locus
HMSN type I CMT1A ² CMT1B CMT1C CMT4A CMT4A CMT4C CMT4C CMTX1 CMTX1 CMTX2 CMTX4	AD/17p11.2-12 ³ AD/1q22-23 ⁴ AD/7 AR/8q AR/11q23 AR/1q XD/Xq13.1 ⁵ XR/Xq22.2 XR/Xq26
HMSN type II CMT2A CMT2B CMT2C CMT2C CMT2D	AD/1p36 AD/3q13-22 AD/? AD/7p14
HMSN type III CMT3A CMT3B CMT3C HNPP	AD/17p11.2-12 ³ AD/1q22-23 ⁴ AD/8q AD/17p11.2-12 ³

(Mendell, 1998; Schaumburg et al., 1992; Schöls, 1998)

AD = autosomal dominant, AR = autosomal recessive, XR = X-linked recessive; XD = X-linked dominant.
 CMT = Charcot-Marie-Tooth. 3 Gene: PMP22 (PMP e peripheral myelin protein). 4 Gene: P₀ (point mutation).
 5 Gene: connexin 32 (point mutation).

396

Table 63	Features of rare nonmetabolic neuropathies (p. 332)
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Syndrome	Inheritance/ Gene Locus	Clinical Features/NCV ¹
Giant axon neuropathy (GAN)	AR/16q24.1	PNP syndrome, fair, frizzy hair, gait impairment, NCV slightly $\ensuremath{\mathbb{I}}$
HSN ² type I	AD/9q22.1-22.3	Sensory and autonomic neuropathy, reflexes [], sensory loss in feet, restless legs, perforating foot ulcers, hearing loss, lancinating pain in limbs, deforming arthropathy, normal motor NLV
FAP ³	AD	Autonomic PNP (⇔ autonomic dysfunction), dissociated sensory loss, pain, trophic disturbances, vitreous opacity, cardiomyopathy, nephropathy, hepatopathy

1 Nerve conduction velocity. **2** Hereditary sensory neuropathy. **3** Familial amyloid polyneuropathy (PNP). There are different subtypes with variable serum protein changes (transthyretin, apolipoprotein A1, gelsolin) that give rise to extracellular amyloid (AF) deposits. Liver transplantation can be performed to remove the amyloid precursors and bring about degeneration of the amyloid deposits.

Table 64 Myopathy syndromes (p. 334)

Features	Potential causes
Acute generalized weakness	Myasthenia gravis, botulism, periodic paralysis ¹ , polymyositis/dermatomyositis, acute rhabdomyolysis, critical illness myopathy, toxic or drug-induced myopathy ² , hypermag- nesemia
Subacute or chronic, mainly pro- ximal weakness	Myasthenia gravis, Lambert–Eaton syndrome, muscular dystrophy, congenital myopathy, polymyositis/dermatomyositis, metabolic myopathy, mitochondriopathy, electrolyte imbalance, endocrine disorder ³ , toxic or drug-induced myopathy
Subacute or chronic, mainly dis- tal weakness	Inclusion body myositis, myotonic dystrophy, facioscapulohumeral muscular dystrophy, nemaline myopathy, central core disease, scapuloperoneal syndrome, Welander myo- pathy ⁴ , oculopharyngodistal myopathy
Periodic weakness	Myasthenia gravis, Lambert-Eaton syndrome, dyskalemic paralysis, paramyotonia con- genita, neuromyotonia, Conn syndrome, thyrotoxicosis
Asymmetric or lo- calized weakness	Facioscapulohumeral muscular dystrophy, myasthenia gravis, ischemic muscular necro- sis, local myositis, muscle rupture/trauma
Multiple system in- volvement	Mitochondriopathy, critical illness myopathy, myotonic dystrophy, proximal myotonic myopathy, dermatomyositis
Dysphagia	Myasthenia gravis, polymyositis, myotonic dystrophy, oculopharyngeal muscular dystro- phy, inclusion body myositis, mitochondriopathy
Myalgia	Viral/bacterial/parasitic/granulomatous/interstitial myositis, dermatomyositis/ polymyosi- tis, vasculitis, eosinophilic fasciitis, polymyalgia rheumatica, fibromyalgia. Alcohol, drugs, hypothyroidism. Metabolic myopathies. Muscle strain. Neuromyotonia. Stiff-man syndrome
Muscle cramps	Idiopathic, exercise-induced, pregnancy, uremia, hypothyroidism, electrolyte imbalance
Muscle hypertrophy	Muscular dystrophy (Duchenne/Becker: calves, deltoid), myotonia congenita, amyloido- sis, cysticercosis, acromegaly, glycogen storage disease type II (Pompe)
Cardiomyopathy	Duchenne/Becker muscular dystrophy, Emery–Dreifuss muscular dystrophy, myotonic dystrophy, centronuclear myopathy, nemaline myopathy, glycogen storage disease type II
CK-emia⁵	Physical stress, muscle trauma (fall, injection, epileptic seizure), toxins/drugs (alcohol), hypothyroidism, female carrier of trait (Duchenne/Becker muscular dystrophy), incipient myopathy (muscular dystrophy, myositis, glycogen storage disease), hereditary

1 Hypokalemic or hyperkalemic form. 2 Table 66. 3 Hypothyroidism or hyperthyroidism, acromegaly, Cushing disease, hyperparathyroidism, Conn syndrome, Cushing syndrome. 4 Myopathia distalis tarda hereditaria. 5 Elevation of creatine kinase (CK) levels in serum without clinical evidence of myopathy; risk of malignant hyperthermia (related to general anesthesia, see p. 346).

Туре	Myopathy	Inh.1	Gene locus	Gene product	
Dystrophinopathy	Duchenne MD ² Becker MD	XR ³ XR	Xp21.2 Xp21.2	Dystrophin Dystrophin	
Sarcoglycanopathy Other LGMDs Other MDs	LGMD2D ⁴ LGMD2E LGMD2C LGMD2F LGMD2B LGMD1A LGMD1B LGMD1C Facioscapulohumeral MD Oculopharyngeal MD Myotonic MD ⁸ Emery-Dreifuss MD	AR ⁵ AR AR AR AR AD AD AD AD AD AD XR	17q21 4q12 5q33 15q15.1-21.1 2p13.3-13.1 5q31 1q11-21 3p25 4q35 14q11.2-q13 19q13 Xq28	α-Sarcoglycan ⁶ β-Sarcoglycan δ-Sarcoglycan Calpain-3 Dysferlin Myotilin Lamin A/C Caveolin-3 ? PABP27 DMPK ⁹ Emerin	
Channel diseases Chloride channel Sodium channel Calcium channel	Thomsen MC ¹⁰ Becker MC Hyperkalemic PP ¹¹ Paramyotonia congenita PAM ¹² Hypokalemic PP ¹³ Malignant hyperthermia Malignant hyperthermia Central core disease	AD AR AD AD AD AD AD AD AD	7q35 7q35 17q23.1-25.3 17q23.1-25.3 17q23.1-25.3 1q32 1q31-32 19q13.1 19q13.1	Chloride channel Chloride channel Sodium channel Sodium channel Calcium channel ¹⁴ Calcium channel ¹⁵ Calcium channel ¹⁵	
Mitochondriopathies	Mitochondrial myopathy ¹⁶		Mitochondrial DNA ¹⁷		

Table 65 Some hereditary myopathies (p. 334)

(Gene loci as specified by OMIM)

Mode of inheritance. 2 Muscular dystrophy. 3 X-linked recessive. 4 LGMD = limb girdle muscular dystrophy.
 5 Autosomal recessive. 6 Adhalin-7-poly(A) binding protein-2. 7 Poly(A) binding protein-2. 8 Unstable trinucleotide repeat (CTG). 9 Myotonin-protein kinase. 10 Myotonia congenita. 11. Hyperkalemic periodic paralysis.
 12. Potassium-sensitive myotonia (myotonia fluctuans). 13. Hypokalemic periodic paralysis.
 14. Dihydropyridine receptor. 16 Mainly systemic diseases, usually maternally inherited or sporadic. 17. Nuclear DNA mutations are rare.

Table 66 Some acquired myopathies (p. 334)

Туре	Myopathy
Neuromuscular end plate dysfunction	Myasthenia gravis, Lambert–Eaton syndrome, botulism
Endocrine myopathy	Hyperthyroidism, hypothyroidism, Cushing syndrome, acromegaly, Conn syn- drome, primary hyperparathyroidism
Inflammatory myopathy	Polymyositis, dermatomyositis, myositis with vasculitis, Churg–Strauss syndrome, granulomatous myositis, inclusion body myositis, myositis induced by pathogens (e.g., bacteria, viruses, parasites)
Toxic/drug-induced myopathy ¹	Alcohol, corticosteroids, lovastatin, simvastatin, cocaine, emetin, diazocholesterol

1 Rarely caused by other drugs or toxins (not listed).

Table 67	Additional	diagnostic	studies fo	or myopathy	(p. 334)
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Method	Information Supplied/Parameters
Pharmacological tests	Edrophonium chloride (p. 404) <i>In vitr</i> o testing for malignant hyperthermia.
Neurography/Stimulation electromyography	Used to exclude peripheral neuropathy (p. 404). Serial stimulation: Evidence of neuromuscular conduction disturbances
Needle electromyo- graphy ¹	 Muscular dystrophy: Possible findings include fibrillation, positive waves, pseudomyotonic discharges. Brief, low-amplitude MAPs, polyphasia rate û, rapid and dense interference pattern Myositis: Fibrillation, positive waves, pseudomyotonic discharges. Polyphasia rate û, narrow and low-amplitude MAPs Myotonic/paramyotonia: Myotonic discharges, MAPs resembling those of muscular dystrophy
Laboratory tests	 Creatine kinase²: > 10 000 in acute rhabdomyolysis, myositis, toxic myopathy, Duchenne/Becker muscular dystrophy (early stage) 4000-10 000 in Duchenne/Becker type muscular dystrophy (later stages), myositis 1000-4000 in muscular dystrophies, hypokalemic or hypothyroid myopathy, congenital myopathy, female carrier (muscular dystrophy) ≤ 1000 in spinal muscular atrophy, amyotrophic lateral sclerosis, inclusion body myositis, chronic/infectious myositis <i>Myoglobin:</i> Severe muscle degeneration ⇒ myoglobinuria³ Serum lactate/pyruvate (venous): Elevated at rest or after light physical exercise ⇒ mitochondriopathies, respiratory chain defects. Absence of rise in disorders of glycolysis and glycogenolysis⁴ <i>Molecular genetics:</i> Depends on results of immunohistochemistry (dystrophies) and biochemical muscle analysis (mitochondriopathies). Used to supplement clinical findings if necessary (channel diseases) Other tests⁵: Erythrocyte sedimentation rate, hepatitis antigen, ANCA⁶ (vasculitis). Eosinophilia (eosinophil fascitits, churg-Strauss syndrome). Sarcoplasmi en PROMM?). Basal TSH. Rheumatoid factor (myositis). Antibodies: AChR⁸ (myasthenia), Jo-1 (myositis, antisynthetase syndrome), Pm-ScI (myositis with systemic sclerosis). SS-A (Ro) (myositis with Sjögren syndrome), U₁RNP (mixed connective tissue disease)
Diagnostic imaging	Ultrasound, CT/MRI: Distribution of atrophy, fat and connective tissue. Supportive evidence when selecting site of biopsy. Localization of local muscle changes (tumor, hemorrhage, pyomyositis/ossification \Rightarrow scintigraphy)
Muscle biopsy	Mainly used for definitive proof of an inflammatory, vasculitic or metabolic my- opathy. Also used for clarification of diseases not clearly classifiable as "myo- genic" or "neurogenic." Sporadic cases of muscular dystrophy

 For abbreviations, see p. 391. 2 U/I of CK-MM; selected examples. 3 Alcohol, barbiturates, acute myositis, malignant hyperthermia, carnitine palmitoyl transferase deficiency, glycogen storage disease type V/VII, posttraumatic, postictal, idiopathic. 4 Determined by stress testing (ischemia = risk of rhabdomyolysis). 5 Selected examples, see also p. 391. 6 Antineutrophil antibodies. 7 Proximal myotonic myopathy. 8 Acetylcholine receptor.
 9 Specimens taken from muscle moderately affected by disease process. Two specimens are deep-frozen in an isopentane–nitrogen mixture (for histochemistry, immunohistochemistry; biochemical diagnosis, etc.). One specimen is fixed in glutaraldehyde for electron-microscopic study.

Criteria	Duchenne MD ²	Becker MD	Limb girdle MD	Facioscapulo- humeral MD	Myotonic MD
Mean age at onset (years)	2	12	Adolescence/ adulthood	Adolescence/ adulthood	Adolescence/ adulthood
Sex ³	М	М	M/F	M/F	M/F
Site of onset	Pelvic girdle	Pelvic girdle	Pelvic girdle (often)	Face, shoulder girdle	Head, shoulder girdle, arms, dorsiflexors of foot
Facial muscle involvement	No	No	No	Yes	Yes (cataract)
(Pseudo) Hy- pertrophy	Calf, deltoid, gluteal muscles	Calf muscles	Calf muscles (rarely)	None	None
Cardiac involve- ment	Common	Rare	Occasional	None	Common (pacemaker)
Inheritance	X-linked reces- sive	X-linked reces- sive	Usually auto- somal recessive	Autosomal dominant	Autosomal dominant
CK level ⁴	50 (to 300) times higher than normal	20 (to 200) times higher than normal	< 10 times higher than normal	Normal to 4 times higher than normal	Normal to 3 times higher than normal
Dystrophin	Absent	Deficient	Normal	Normal	Normal
Myotonia	No	No	No	No	Yes
Prognosis	Age (years) 3–6, gait dis- turbance; 5–6, hypertrophy; 6–11, increas- ing weakness and contrac- tures; death often at age 15–30	Slow progres- sion. Unable to walk by ca. age 20. Mean age at death, 42 years (range, 23–89)	Mainly slow. Life span usu- ally only slightly decreased	Slow progres- sion. Ability to walk is pre- served. Normal life span	Ability to walk is preserved. Life span short- ened only in severe cases

Table 68 Clinical features of selected muscular dystrophies¹ (p. 336)

1 Very rare syndromes (prevalence < $1/10^6$) such as Emery–Dreifuss MD, oculopharyngeal MD, distal myopathies, proximal myotonic MD, and congenital MD are not listed here. 2 MD = muscular dystrophy. 3 M = male; F = female. 4 CK = creatine kinase.

Criterion	Hyperka- lemic par- alysis	Paramyo- tonia con- genita	Myotonia fluctuans	Myotonia congenita Thomsen	Myotonia congenita Becker	Proximal myotonic myo- pathy ¹	Myotonic dystrophy
Periodic paralysis	Yes	Yes	No	No	Yes	No	No
Cold-in- duced par- alysis	No	Yes	No	No	No	No	No
Potassium- induced paralysis	Yes	Some- times	No	No	No	No	No
Paradoxi- cal myo- tonia ²	Some- times	Yes	No	No	No	No	No
Additional organ in- volve- ment ³	No	No	No	No	No	Yes	Yes
Inheri- tance	AD ⁴	AD	AD	AD	AR ⁵	AD	AD
Defect	Sodium channel	Sodium channel	Sodium channel	Chloride channel	Chloride channel	?	Protein kinase
Features	Duration of para- lytic at- tacks var- ies (≤ 4 hours)	Attacks of muscle stiffness and weak- ness can last up to 1 day	Myotonia of variable severity	General- ized myo- tonia, no weakness	Myotonia more severe than in Thomsen type. Tran- sient weakness	Proximal muscle weakness, cataract, muscle pain, mild muscular atrophy	Weakness, especially of craniocer- vical muscles, less pro- nounced in limbs (mainly distal). Cataract. Defective cardiac impulse conduc- tion

Table 69 Myotonia and periodic paralysis (p. 338)

(Ptacek et al., 1993)

1 Unlike in myotonic dystrophy, there is no cytosine-thymine-guanine (CTG) repeat. 2 In this case, myotonia generally subsides after repeated voluntary muscle contraction ("warm-up"), whereas it increases in paradoxical myotonia. 3 I.e., extramuscular involvement. 4 Autosomal dominant. 5 Autosomal recessive.

Table 70	Selected	forms o	f congenital	myopathy (p. 340))
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Myopathy	Gene Locus/Gene	Comments
Central core disease	AD1: 19q13.1/Ryanodine receptor	Neonatal hypotonia. Slow progression. Atten- tuated muscles, skeletal anomalies ² , hyporeflexia or areflexia; exercise-induced muscle stiffness; risk of malignant hyperthermia
Nemaline myopathy (NEM1)	AD: 1q22–23/Tropomyosin-3	Neonatal hypotonia; nonprogressive; high palate
Nemaline myopathy (NEM2)	AR3/2q22/Nebulin	Delayed motor development. Attenuated muscles, thin extremities; dysplasia ⁴ . Respiratory disturbances (paretic diaphragm muscles), recur- rent pneumonia, dysphagia, dysarthria; hy- poreflexia or areflexia
Centronuclear (my- otubular) myopathy	XR ⁵ : Xq28/Myotubularin	Neonatal hypotonia, facial muscle weakness, ex- ternal ophthalmoplegia, hyporeflexia, respiratory disturbances, dysphagia, high palate

1 Autosomal dominant. 2 Congenital hip dislocation, chest deformity, kyphoscoliosis, pes cavus. 3 Autosomal recessive. 4 Elongated and oval face, open mouth, micrognathia, high palate, kyphosis, hyperlordosis, pes cavus, cardiomyopathy, heart failure. 5 X-linked recessive; autosomal recessive and autosomal dominant inheritance are also found, with less severe manifestations.

Appendix

Table 71 Metabolic myopathies (p. 340)

Myopathy/Gene Locus	Defect/Inheritance	Features
Carbohydrate metabolism • Acid maltase deficiency ¹ (type II, Pompe)/17q25.2–25.3 • Muscle phosphorylase deficien-	 1,4-Glucosidase/ Autosomal recessive Myophosphorylase/ 	 Slowly progressive proximal myopathy, respiratory distur- bances (nocturnal hypoventila- tion) Exercise-induced muscle pain
cy ¹ (type V, McArdle)/11q13	 Autosomal recessive 	and stiffness, contractures that subside at rest, rhabdomyolysis
 Phosphofructokinase deficiency (type VII, Tarui)/12q13.3 	 Phosphofructokinase/ Autosomal recessive	Similar to McArdle type
 Fat metabolism Carnitine deficiency my- opathy/? CPT-I deficiency²/11q13 	Carnitine/Autosomal recessive?CPT I/Autosomal recessive	 Symmetrical, proximal, slowly progressive myopathy, CK 0 Exercise and cold-induced muscle pain and weakness, rhabdomyolysis, hypoqlycemia,
CPT-II deficiency/1p32	CPT II/Autosomal recessive	 byperammonemia Exercise/fasting-induced muscle pain, rhabdomyolysis

(continued next page)

Table 71	Metabolic	myopathies	(p. 340)	(continued)
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Myopathy/Gene Locus	Defect/Inheritance	Features
Mitochondria • CPEO ³	• mtDNA deletion in ca. 50% of cases	 Ptosis, external ophthalmople- gia, tapetoretinal degenera- tion, cardiac arrhythmias, pro-
• KSS ⁴	mtDNA deletion/duplication	 ximal myopathy Onset before 13th year of life, ataxia, hearing impairment,
 MERRF⁵ MELAS⁶ 	 mtDNA point mutation mtDNA point mutation 	 Myoclonus, ataxia, seizures Episodic vomiting, focal seizures, dwarfism, proximal muscle weakness
 LHON⁷ 	mtDNA point mutation	 Acute/subacute bilateral loss of vision, eye pain
• MILS ⁸	 mtDNA point mutation⁹ 	 Developmental delay, ataxia, dystonia, visual disturbances, respiratory disturbances¹⁰

1 Adult type. 2 Carnitine palmitoyl transferase; defect located on outer mitochondrial membrane in type I, and on inner membrane in type II. 3 Chronic progressive external ophthalmoplegia. 4 Kearns–Sayre syndrome; cardiac pacemaker implantation may be necessary in patients with cardiac arrhythmias. 5 Myoclonus epilepsy with ragged red fibers. 6 Myopathy, encephalopathy, lactic acidosis, and "strokelike episodes". 7 Hereditary hepaticoptic neuropathy. 8 Maternally inherited Leigh syndrome. 9 Autosomal recessive and sporadic forms are also found. 10 T2-weighted MRI reveals bilateral symmetric lesions (brain stem, periaqueductal region, cerebellum, basal ganglia)

Alternatives
Cephalosporins, chloramphenicol
Promethazine, thioridazine. Chlordiazepoxide, ma- protiline, mianserin or carbamazepine can be used at low doses and with careful monitoring
Carbamazepine
Digitalis, reserpine, methyldopa, tocainide, verapamil (low-dose)
Aspirin, gold, indometacin, acetaminophen, di- clofenac, local/regional anesthesia, spinal anesthesia, inhalant anesthetics/deeper general anesthesia

Table 72	Drugs that can	aggravate m	vasthenia	aravis (p. 342)

1 Mask symptoms of myasthenia. 2 High starting dose.

(Selected drugs from McNamara and Guay, 1997)

Syndrome	Symptoms and Signs	Precipitating Factors
Myasthenic crisis	Restlessness, anxiety, confusion, respira- tory weakness, weak cough, dysphagia, dysarthria, mydriasis, ptosis, tachycardia, pallor	Infectious diseases, surgical interventions, anesthesia, drugs, psychosocial stress, im- paired drug uptake (vomiting, diarrhea), disease progression, previously unde- tected myasthenia (and previously men- tioned factors)
Cholinergic crisis	Restless, anxiety, confusion, respiratory weakness, weak cough, dysphagia, dy- sarthria, miosis, bradycardia, skin redden- ing, muscle fasciculation/spasms, saliva- tion, tenesmus, diarrhea	Overdosage (relative) of AChE inhibitors; acetylcholine poisoning

Table 73 Myasthenia-related crises (p. 342)

Table 74 Ancillary tests in myasthenia gravis (p. 342)

Test	Objective	Interpretation of Results
Edrophonium chloride test ¹ (Ten- silon, Camsilon)	Increase in muscle strength (with improvement of ptosis, eye move- ments, speech, and swallowing)	Marked improvement (beginning 30 seconds after administration and lasting roughly 5 minutes) ⇒ unequivocal response. Sensitivity for OMC ² : ca. 86%, for GMG ³ : ca. 95%
Electromyography (EMG) ⁴	Documentation of impaired neu- romuscular conduction (decre- ment in amplitude seen with se- rial stimulation; jitter may be ob- served in single-fiber EMG)	A decrement of 10% or more is pathological. Sensitivity of serial stimulation in OMG: ca. 34%; in GMG: up to 77%. Prior muscle ex- ercise \Rightarrow more pronounced decre- ment. Sensitivity of single-fiber EMG: ca. 92%
Serum acetylcholine receptor anti- body titer	Documentation of presence of acetylcholine receptor antibodies	Sensitivity: 50% in OMG, ca. 90% in GMG. False-positive results may occur in Lambert-Eaton syn- drome, rarely in amyotrophic lateral sclerosis
Diagnostic imaging ⁵	Measurement of thymus	Thymic enlargement due to thy- moma or hyperplasia

(Phillips and Melnick, 1990)

1 Short-term inhibition of cholinesterase, given intravenously for diagnostic purposes. 2 Ocular myasthenia gravis. 3 Generalized myasthenia gravis. 4 Example: Repeated stimulation of accessory nerve (3/sec for 3 seconds) and recording of activity in trapezius muscle. 5 CT (contrast-enhanced) or MRI (younger patients, better differentiation of thymic hyperplasia).

Table 75 Toxic myopathies (p. 347)

Syndrome	Substances (selected)
Muscle weakness with or without pain; rhabdomyo- lysis may occur	Alcohol, chloroquine, cimetidine, clofibrate, cocaine, colchicine, ciclosporin, disulfiram, emetine, er- gotamine, gemfibrozil, induced hypokalemia (diuret- ics, licorice), imipramine, isoniazide, lithium, lovastatin, meprobamate, niacin, pentazocine, thy- roid hormones, vincristine, zidovudine
Myalgia	Alcohol, allopurinol, cimetidine, clofibrate, clonidine, dihydroergotamine, ergotamine, methyldopa, suc- cinylcholine, vincristine, zidovudine
Polymyositis, pseudo-lupus erythematosus	Bezafibrate, chlorpromazine, cimetidine, clofibrate, D-penicillamine, etofibrate, etofyllin clofibrate, fenofi- brate, gold, hydralazine, isoniazide, L-tryptophan, penicillin, phenytoin, procainamide, tetracyclines, zidovudine
Myotonia	Ciclosporin, 20,25-diazocholesterol, diuretics, D-peni- cillamine, fenoterol, pindolol, propranolol
Local muscle lesions (pain, swelling, local muscular atrophy)	Heroin, insulin, meperidine, pentazocine

Site of Lesion	Syndrome ⇒ Manifestation	Symptoms and Signs	Common Tumors	Lesions/ Antibodies
Motor neuron	Subacute muscular atrophy (hands, bulbar muscles) ⇔ weeks to months	Asymmetrical para- lysis, muscle atro- phy (p. 304)	Small-cell lung cancer, lymphoma, renal cell carci- noma	Motor neurons/ Anti-Hu ¹
Spinal posterior root, ganglion	Subacute sensory neuronopathy ⇔ weeks to months	Marked sensory loss, areflexia, ataxia, paresthe- siae, pain	Small-cell lung cancer, other lung tumors	Spinal ganglia
Proximal peripheral nerve	 Acute poly- radiculopathy ⇒ hours to days Chronic poly- radiculopathy³ ⇒ weeks to months 	 Ascending sensorimotor deficits² Chronic progressive/recurrent sensorimotor deficits 	 Hodgkin disease Small-cell lung cancer, lym- phoma, my- eloma 	Segmental demy- elination, neuritis
Distal peripheral nerve	 Paraproteinemic polyneuropathy ⇒ weeks to months Sensorimotor polyneuropathy ⇒ weeks to months Neuromyotonia 	 See p. 328 Distal symmetrical polyneuropathy Muscle stiffness, cramps 	 Plasmacytoma Small-cell lung cancer, other cancers Thymoma, lung cancer 	 Segmental de- myelination Mainly axonal lesions Distal motor nerve/Anti- VGPC⁴
End-plate region	 Lambert-Eaton syndrome ⇒ weeks to months Myasthenia gravis ⇒ weeks to months 	See p. 342See p. 342	 Small-cell lung cancer; breast, prostate or stomach cancer Thymoma 	 See p. 343/ Anti-VGCC anti- bodies⁵ See p. 343/ Skeletal muscle antibodies
Skeletal muscle	 Polymyositis/ dermatomyosi- tis ⇒ months to years Rhabdomyolysis ⇒ days to weeks 	 See p. 344 Rapidly progressive paralysis, dysphagia 	 Various cancers (breast, lung or ovarian cancer, lymphoma) Various cancers 	 Myonecrosis, lymphomono- cytic infiltrates Myonecrosis, rare inflam- matory infil- trates

 Table 76
 Neuromuscular paraneoplastic syndromes (pp. 347, 388)

(Brown, 1998)

1 In small-cell lung cancer. 2 Similar to Guillain–Barré syndrome (p. 326). 3 Similar to CIDP (p. 328). 4 VGPC = voltage-gated potassium channel; EMG shows high-frequency discharges (150–300 Hz). 5 VGCC = voltage-gated calcium channel.

Table 77 Laboratory tests (p. 351)

Test/Objective	Risks	Comments
 Antiepileptic drugs Verify drug compliance Assess for drug resistance Avoid underdosage or over- dosage Assess for drug interactions 	 Laboratory error Misuse of measured values (the physician should be guided by the clinical objective of a seizure-free state, rather than by "therapeutic levels") 	Time of sample collection is deter- mined by the pharmacokinetics of the antiepileptic drug in question
Lumbar puncture • Measure CSF pressure • Obtain CSF sample for analysis • Intrathecal drug administration • Diagnosis (contrast agent ¹ , radioactive substances ²)	 Increased intracranial pressure³ Intraspinal mass⁴ Postpuncture headache Intraspinal hemorrhage (coagulopathy) Meningitis Discitis 	Suboccipital or lateral cervical puncture is very rarely indicated (e.g. if a CSF sample is required, but cannot be obtained by lumbar puncture, or for myelography above a spinal lesion). Myelogra- phy and MRI have rendered Queckenstedt's test'o obsolete

1 For myelography. 2 For scintigraphy. 3 Risk of transtentorial/cerebellar herniation. 4 Risk of acute spinal decompensation with paraplegia. 5 Compression of jugular vein to test for patency of subarachnoid space, which may be blocked, for example, by a spinal tumor.

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410

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412

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Index

A

Aachen aphasia test 124 Abasia 276 Abetalipoproteinemia 280, 281, 300, 307, 332 Abiotrophy 296 Abscess brain 222, 226, 227 diagnosis 226 pathogenesis 226 candida 248 epidural 222 tuberculous 232 Absidia 248 Abulia 122, 123 Acalculia 128, 129 Acanthocytes 300, 301 Acetazolamide 338 Acetylcholine 140, 152, 210 Acetylcholine receptor antibody titer 404 Acetylcholinesterase inhibitors 298, 342 Acid maltase deficiency 340, 402 Acidosis 162 Acoustic meatus, external 100 Acoustic neuroma 258, 259, 294 Acquired immunodeficiency syndrome (AIDS) 240 cytomegalovirus and 244 progressive multifocal leukoencephalopathy and 244 Acrodermatitis chronica atrophicans 228 ACTH-secreting tumors 258 Acute demyelinating encephalomyelitis (ADEM) 234 Acute dystonic reactions 66, 204, 205 Acute inflammatory demyelinating polyradiculoneuropathy (AIDP) 395 Acute motor-axonal neuropathy (AMAN) 395 Acute motor-sensory axonal neuropathy (AMSAN) 395 Acyclovir 236, 238

Adaptation, olfactory 76 Adenohypophysis 142 Adenoma, pituitary 258, 259, 377 Adie syndrome 92 Adrenal medulla 140 Adrenoleukodystrophy 307 Adrenomyeloneuropathy 332. 384 Ageusia 78 Aging 296, 382 degenerative changes 296 disease and 296 Agnosia 132 body-image 132 finger 132 Agrammatism 124, 126 Agraphia 128, 129 alexia with 128 aphasic 128 apraxic 128 isolated 128 spatial 128, 132 Agyria 381 AIDS see Acquired immunodeficiency syndrome Akathisia 66 Akinesia 206, 208 Huntington disease 300 Akinetic mutism 120, 122, 368 Albendazole 250 Alcohol intoxication 312, 313 withdrawal syndrome 312, 313 Alcoholism 312, 313, 366 fetal alcohol syndrome 314 late complications 314 Alexia 128 agraphia and 128 anterior 128 central 128 isolated 128 Alien hand syndrome 24, 302 Alkalosis 162 Alleles 288 Allodynia 316, 346 Alzheimer disease (AD) 136, 296-298, 299, 366

agraphia 128 pathogenesis 297-298 risk factors 296 symptoms and signs 297 treatment 298 see also Dementia Amaurosis fugax 82, 168, 372 Amblyopia, tobacco-alcohol 314 Amimia 362 Amnesia 134, 268, 365, 368 anterograde 134 examination 134 retrograde 134 Amoxicillin 228 Amphetamine abuse 314 Amphotericin B 248 Ampullary crests 56 Amygdala 144 Amyloid precursor protein (APP) 297 Amvloid-Aß 297 Amyotrophic lateral sclerosis (ALS) 304, 386 adult-onset 304 bladder dysfunction and 371 iuvenile 304 sporadic 304 Amyotrophy, neuralgic 321, 328.329 Anal incontinence 370 Anencephaly 292 Anesthesia 106 Aneurysm 178, 179 fusiform 178 rupture 176, 178, 179 treatment 178 saccular 178 septic-embolic 178, 226 Angiitis cerebral 180 von Heubner 230 Angiography 354 Angiomatosis cutaneous 294 encephalofacial 294 Angiopathy, amyloid 178 Anhidrosis, generalized 152 Anisocoria 90 Annulus fibrosus 30

Anomia 124 Anosmia 76 partial 76 Anosognosia 132 Anterocollis 64 Antibiotics 224 adverse effects 389 myasthenia gravis aggravation 403 see also specific drugs and infections Anticholinergic agents 212 Anticipation 300 Anticoagulants, stroke management 174 Anticonvulsants 198 myasthenia gravis aggravation 403 Antidepressants, adverse effects 389 Antiemetics, acute dystonic reaction 204 Antiepileptic drugs (AEDs) 198, 264 laboratory tests 407 myasthenia gravis aggravation 403 Antimicrobial therapy see Antibiotics Antioxidants, neuroprotective therapy 212 Antiplatelet therapy, stroke 174 Anton syndrome 132 Anxiety, Parkinson disease and 206 Apallic syndrome 117, 120, 121 Aphasia 124, 126-127 Alzheimer disease and 297 amnestic (anomic) 126 Broca's 126, 127 conduction 126 crossed 126 global 126, 127 subcortical 126 test of 124 transcortical 126, 127 motor 126 sensory 126 Wernicke's 126, 127 Aphemia 124 Apnea test 364 Apo E gene 297 Apomorphine 212 Apraxia 128, 129 Alzheimer disease and 297 buccofacial 128

constructional 132 dressing 128, 129, 132 gait 128, 374 ideational 128 ideomotor 128, 129 lid-opening 128, 129, 362 limb 128 Apraxia-like syndromes 128 Aqueduct, cerebral (of Sylvius) 8 Arboviruses 376 Arch aortic 148 150 Archeocerebellum 54 Area Broca's 124 postrema 140 Wernicke's 124 Argyll-Robertson pupils 92, 230 Arousal disorders 116, 117 Arrhythmias, neurogenic 148 Arteriovenous malformations (AVMs) 178, 179 hemorrhage treatment 178 Arteritis 226, 227 Takavasu 180 temporal 180 see also Vasculitis Artery(ies) age-related changes 382 basilar 14-15 occlusion 170, 171 callosomarginal 12 carotid 10-12 common 10 occlusion 168 external 10 internal 10-12 infarction 168 169 occlusion 168 central posterolateral 16 posteromedial 16 cerebellar infarction 170 inferior anterior 14, 170 posterior 14, 170, 171 occlusion 170 superior 14, 170 cerebral anterior 12, 13 infarction 168, 169 middle 12, 13 occlusion 168, 169 posterior 16-17 occlusion 170, 171

pars circularis 16 pars terminalis 16 choroidal, anterior 10 infarction 168 communicating anterior 12 posterior 10, 16 frontobasal 12 frontobasilar 12 frontopolar 12 insular 12 lenticulostriate 12 occlusion 168 medullary 22 meningeal 6 middle 10 occipital lateral 16 medial 16 ophthalmic 10 occlusion 168 paracentral 12 parietal anterior 12 posterior 12 parieto-occipital 12 pericallosal 12 precuneal 12 radicular, great (of Adamkiewicz) 22 occlusion 282 recurrent, of Heubner 12 retinal, central 10 segmental 22 spinal 22, 23, 283 anterior 14, 22 syndrome of 282 posterior 14, 22 syndrome of 282 subclavian occlusion 170 subclavian steal 170 sulcocommissural artery syndrome 282 temporal 12 thalmostriate 12 vertebral 14-15 extracranial 14 intracranial 14 occlusion 170 Arthralgia, postpolio syndrome 242 Articulation 130 dvsarthria 130 Aspergillus fumigatus (aspergillosis) 248, 249 Aspiration 102

416

Aspirin, adverse effects 389 Astasia 276 Asterixis 68, 69 Astrocytoma 256, 377 anaplastic 260, 261, 377 low-grade 256 pilocytic 256, 377 Ataxia 107, 276, 374 autosomal dominant cerebellar (ADCA) 280 episodic (EA) 280 Friedreich (FA) 280, 281 gait 276, 277 idiopathic cerebellar (IDCA) 276 postural 276 spinal (sensory) 276, 374 spinocerebellar (SCA) 280. 384 truncal 276 Ataxia-telangiectasia 280, 295 Athetosis 383 Atlas 30 Atrophy age-related 382 cerebellar 279 alcoholism and 314 cerebral alcoholism and 314 Huntington disease 300 muscular 49-52, 107, 281, 287, 334, 406 peripheral neuropathy and 316 poliomyelitis and 242, 243 spinal (SMA) 385 spinobulbar 385 olivopontocerebellar 302 optic nerve 158, 159 temporal papillary 215 Attack(s) drop 204, 205, 374 panic 202, 203 transient ischemic (TIA) 166 crescendo 166 Attention deficits 122, 123 directed attention 122 divided attention 122 evaluation 353 Auditory evoked potentials (AEP) 218, 352 Auditory pathway 100, 101 Aura migraine 184, 185 seizure 192 Automatism 124, 126

Autonomic dysfunction 48, 371 diabetic neuropathy 324 multiple sclerosis 216, 217 neurosyphilis 230 Parkinson disease 208, 209 peripheral neuropathies 316. 390 Autonomic nervous system (ANS) 2, 140-141 central portion 140, 141 afferent connections 140 efferent connections 140 neurotransmitters 140 enteric 154 peripheral portion 140-141, 144-146 afferent connections 140 efferent connections 140 neurotransmitters 140-141 spinal nuclei 140 Autotopagnosia 132 Axis 30 Axonopathy 316 Axonotmesis 330, 331 Axons 2 Azathioprine 180, 220, 342, 344

B

Babinski sign 40, 49 Baclofen, adverse effects 389 Bacterial infections 226-233 brain abscess 226, 227 Lyme disease 228-229 meningitis/meningoencephalitis 226, 376 neurosyphilis 230–231 septic encephalopathy 226 vasculitis 226 ventriculitis 226, 227 see also specific infections Ballism 66, 383 Bannwarth syndrome 228 Bárány's pointing test 276 Baroreceptors 148 Basilar impression 292, 293 Bassen-Kornzweig syndrome 300 Becker muscular dystrophy 336, 337, 400 Behavioral changes 122-123, 368 brain tumors 254, 255 Huntington disease 300

intracranial hypertension 158 159 multiple sclerosis 216, 217 normal-pressure hydrocephalus 160 Parkinson disease 206-209 stroke 166 Behcet disease 180, 234 Benedict syndrome 70, 358 Benperidol 204 Benzodiazepines 198 Biopsy 354, 399 sural nerve 391 "Black-curtain" phenomenon 168 Bladder dysfunction 156, 371 multiple sclerosis 216, 217. 218 normal-pressure hydrocephalus and 160, 161 Parkinson disease 208 Bladder function 156, 157 tests of 218 residual urine volume 218 urodynamic electromvography 218 Blepharospasm 64, 65, 362 Blindness, transient monocular 168.372 Blink reflex 96, 98 Blood pressure 143, 148, 367 Parkinson disease and 208 see also Hypertension; Hypotension Blood-brain barrier 8-10 disruption of 224 multiple sclerosis and 220 passage of pathogens 224 Blood-CSF barrier 8-10 Body(ies) geniculate, lateral 80 inclusion 52, 252, 344, 345 Lafora 307 Lewy 208, 210, 302 para-aortic 150 Body image perception disturbances 132 Body temperature see Thermoregulation Body-image agnosia 132 Bone windows 4 Borrelia burgdorferi 228–229 Bourneville-Pringle disease 294.295 Bovine spongiform encephalopathy (BSE) 252 Brachycephaly 381

Bradykinesia 206 Huntington disease 300 Bragard's sign 318 Brain 2 abscess 222, 226, 227 diagnosis 226 pathogenesis 226 blood-brain barrier 8-10 degenerative changes 296. 382 fore brain 2 hind brain 2 mid brain 2, 26 syndromes 70, 71, 358-359 traumatic brain injury (TBI) 266-271 complications 269, 270 evaluation 266 pathogenesis 270, 271 primary injury 266, 270 prognosis 268 secondary sequelae 268. 270 treatment 270 hospital 270 scene of accident 270. 271 types of 266 see also Brain tumors Brain death 120 Brain stem 2, 3, 24, 26-27 encephalitis 222 hemorrhage 176 syndromes 70-71 Brain tumors 254-265 ACTH-secreting 258 aging and 296 benign 256-257 bladder dysfunction and 371 classification 264 grades of malignancy 377 growth hormone-secreting 258 incidence 264 infratentorial region 258 malignant 260-261 metastatic disease 262-263 treatment 265 severity 264, 377 supratentorial region 258 symptoms and signs 254–255 behavioral changes 254, 255 epileptic seizures 254 focal neurological signs 254, 255

headache 254, 255 intracranial hypertension 254 nausea, vertigo and malaise 254, 255 treatment 264-265 aftercare 265 grade I tumors 264 grade II tumors 264 grade III tumors 264-265 grade IV tumors 265 metastases 265 symptomatic treatment 264 see also specific tumors Breathing 150, 151 disorders 150, 151 Brivudine 238 Broca's aphasia 126, 127 area 124 Bromocriptine 212 Bromopride 204 Brown-Séquard syndrome 48 Brudzinski's sign 222 Bruxism 114 Bundle, medial fore brain 144 **Bupidine 212** Burns 312 Burst fracture 380

С

Cabergoline 212 Calcium antagonists acute dystonic reaction 204 adverse effects 389 Calcium balance 310 Calcium channel dysfunction 338.398 Caloric testing 26 Calvaria 4 metastases 262 Canal ear 100 infraorbital 4 semicircular 56 spinal 30, 31 vestibular 100 Canalolithiasis 58 Candida albicans (candidosis) 248, 249 Capsule, internal, hemorrhage 176 Carbamazepine 198, 264 Carcinoma choroid plexus 377 meningitis and 262

Cardiomyopathy 397 Carnitine deficiency 340, 402 Carnitine palmitoyl transferase (CPT) deficiency 402 Carpal tunnel syndrome 322 Cataplexy 374 Cataract 382 myotonic 338 Cauda equina 2 syndrome 319 Causalgia 110 Cavernoma 178, 179 Cefotaxime 228 Ceftriaxone 226, 228 Central core disease 402 Central nervous system (CNS) 2 infections 222-225 clinical manifestations 222 course of 222 localization 222, 223 opportunistic infections 240 pathogenesis 224, 225 prophylaxis 224 treatment 224 see also specific infections see also Brain; Spinal cord Central pontine myelinolysis 310. 315 alcoholism and 314 Central salt-wasting syndrome Cerebellitis, acute 238 Cerebellum 2, 24, 42, 54-55 diseases 276-281 acquired 278-279 atrophy 279 alcoholism and 314 diagnostic studies 276 hereditary 280-281 autosomal dominant 280 autosomal recessive 280 signs of dysfunction 276 topography of lesions 276 see also specific diseases hemorrhage 176 Cerebral blood flow (CBF) 162 hemodynamic insufficiency 174 hypoperfusion 174 Cerebral cortex 24-25 Brodmann classification 24, 25 cvtoarchitecture 24 functional areas 24 ischemia 148

418

motor 42 primary 42 lesions 46 supplementary 42 premotor area 42 projection areas 24 Cerebral palsy, infantile 288-291, 381 causes 288 symptoms and signs 288 treatment 290 Cerebral perfusion pressure (CPP) 162 Cerebral vascular resistance (CVR) 162 Cerebral ventricles 8, 9 Cerebritis 222, 226 Cerebrospinal fluid (CSF) 8-9 blood-CSF barrier 8-10 circulation 8, 9 impaired 160, 161, 372 intracranial hypotension and 160 leak 269, 372, 379 lymphoma and 260 multiple sclerosis and 218 pressure measurement 161 viral meningoencephalitis and 234 volume 162 Cervical syndrome 188, 189 upper 188 Charcot joints 230 Chemodetectoma 258 Chemoreceptors 104, 150, 151 Chemotherapy 264-265 adverse effects 389 Cheyne-Stokes respiration 118 Chiari malformation 292, 293 Chiasm, optic 80 lesions 82 Chickenpox 238 symptoms and signs 238 Chloride channel disease 398 Chlorprothixene 264 Cholinergic crisis 404 Chondrosarcoma 260 Chorda tympani lesions 78 Chordoma 258, 259 Chorea 66-67, 300, 383 secondary 66 Svdenham's 66 Choreoathetosis 66 Chromosomal anomalies 381 Chronic inflammatory demyelinating polyradiculoneuropathy (CIDP) 327, 328

Chronic paroxysmal hemicrania (CPH) 186, 190 Chronobiology 112 Chronopathology 112 Churg-Strauss syndrome 180. 344 Cidofovir 244 Cingulate gyrus lesions 122 Cinnarizine 204 Ciprofloxacin 226 Circadian rhythm 112, 113 disturbances 114 Circle of Willis 10, 12, 13 Circulation 148-149 anterior 10 central nervous regulation 148 cerebrospinal fluid 8, 9 impaired 160, 161, 372 posterior 10 Cistern(s) 8 ambient 8 cerebellomedullary (cisterna magna) 8 cerebellopontine 8 chiasmatic 8 interpeduncular 8 posterior 8 Claudication, spinal 282, 284, 285 Clindamycin/folinic acid 250 Clock/numbers test 136, 137 Cluster headache see Headache Cocaine abuse 314 Cochlea 100, 101 Cognitive impairment Alzheimer disease 297 Huntington disease 300 Colon 154 Column Clarke's 104 vertebral 30 Coma 92, 118-119 pupillary dilatation 92 pupilloconstriction 92 staging 118-119, 267 brain stem reflexes 118 Glasgow coma scale 378 respiratory pattern abnormalities 118 spontaneous movement 118 stimuli 118 Comalike syndromes 120-121 akinetic mutism 120, 122, 368 locked-in syndrome 120, 121, 170, 359

persistent vegetative state 120 Commissure, anterior 144 Complex regional pain syndrome (CRPS) 110 Compliance 162 Compression fracture 380 nerve injuries 330 Computed tomography (CT) 354 brain tumors 260, 265 head trauma and 266 multiple sclerosis 218 Confabulation 134 Confusion 116, 117, 368 Alzheimer disease 297 Connective tissue diseases 234 Consciousness 116-117 acute disturbances 116-117 arousal disorders 116, 117 confusion 116, 117 somnolence 116, 117 stupor 116, 117 assessment of 116 content of 116 head trauma and 266, 267 assessment 379 level of 116 normal state of 116, 117 psychogenic disturbances 120 stroke and 166 see also Coma **Constipation 370** Parkinson disease 208 Continence 156 see also Incontinence Conus medullaris 2 Conversion disorders 138 Convulsions, neonatal 196 see also Seizures Coordination dysfunction 276 multiple sclerosis 216, 217 Corneal reflex 26, 96 Corpus callosum 24 agenesis 290 Cortex auditory 100, 101 primary 100 secondary 100 entorhinal 144 premotor, lesions 122 somatosensorv 108 see also Cerebral cortex Corticobasal degeneration (CBD) 302, 303

Corticosteroids 143, 264, 342, 344 adverse effects 389 multiple sclerosis treatment 220 Cortisol 367 Cough reflex 26 Coumarins, adverse effects 380 Craniocervical junction anomalies 292 Craniopharyngioma 258, 259, 377 adamantinomatous 258 papillary 258 treatment 264 Craniostenosis 381 Cranium 4 roof 4 see also Skull Creatine kinase elevation 397 Creutzfeldt-Jakob disease (CID) 252, 253 Crisis cholinergic 404 mvasthenic 404 Critical illness myopathy (CIM) 347.379 Critical illness polyneuropathy (CIP) 347, 379 Crow-Fukase syndrome 328 Cryptococcus neoformans (cryptococcosis) 248, 249 Cupulae 56 Cyanocobalamin 286 Cyclophosphamide 180, 220, 328 Cvst arachnoid 290, 291 colloid 258, 259 porencephalic 290 Cysticercosis 250, 251 Cytokines 220 Cytomegalovirus (CMV) 244-245 pathogenesis 244 symptoms and signs 244

D

420

Dandy–Walker malformation 292 Dantrolene 347 Death 120–121, 364 Debrancher deficiency 340 Decerebration syndrome 46, 47, 118, 158, 159 Decortication syndrome 46, 47, 118 Deep brain stimulation 212 Deficiency acid maltase 340, 402 carnitine 340, 402 carnitine palmitovl transferase (CPT) 402 debrancher 340 folic acid 286 muscle phosphorylase 402 phosphofructokinase 402 vitamin B1 312 vitamin B12 286, 287 vitamin E 280 Degeneration 382 corticobasal (CBD) 302, 303 panthothenate kinaseassociated 307 striatonigral (SND) 302 subacute combined (SCD) 286.287 Degenerative changes 296 radicular syndromes and 392 see also Neurodegenerative diseases Deglutition 102, 103 disturbances 102, 103 mechanism 102 nerve pathways 102, 103 Dehydration 372 Dejerine-Sottas disease 332 Delirium 116, 368 tremens 312 Dementia 136-137 alcoholic 314, 366 bladder dysfunction and 371 classification 366 diagnosis 136 differential 383 dialysis 310 examination 136 frontotemporal (FTD) 298, 299 Parkinson disease and 208 thalamic 116 vascular 298, 299, 366 multi-infarct 298 strategic infarct 298 with Lewy bodies 208, 302 see also Alzheimer disease Demyelination 220 Dens fracture 380 Depersonalization 202 Depression 366 cortical spreading 184

differential diagnosis 383 Parkinson disease and 206 Derealization 202 Dermatomes 32-36 Dermatomyositis 344, 345. 406 Developmental anomalies 288. 381 Diabetes mellitus 324, 325 Diabetic ketoacidosis 308 neuropathy 324, 325, 395 diagnosis 324 symptoms and signs 324 treatment 324 pandysautonomia 395 Dialysis encephalopathy 310, 311 Diaphragma sellae 6 Diarrhea 370 Diencephalon 2 Diffuse Lewy body disease (DLB) 302 Diploë 4 Diplopia 86 multiple sclerosis 214 Disconnection syndrome 24, 128 Disequilibrium syndrome 310 Disk(s) intervertebral 30 herniation 318, 319, 392 optic 80 Dislocation atlantoaxial 380 fracture 380 Disorientation 132-133 right-left 132 "Doll's eye" phenomenon 26, 302.303 coma and 118 Dopamine 210 age-related changes 382 Parkinson disease and 210 Dopamine agonists 208, 212 adverse effects 389 Dopaminergic agents 212 Doxycycline 228 Drooling 206, 207 Drop attacks 204, 205, 374 Drop metastasis 260, 262 Duchenne muscular dystrophy 52, 53, 336, 337, 400 Duplex sonography 353 Dura mater 6 fistula 178, 282, 283 injuries 266 spinal 30

Dysarthria 124, 130, 166, 276, 365 Dysarthrophonia 206, 276 Dysdiadochokinesia 276, 277 Dysesthesia 106, 316 Parkinson disease and 208 Dysgeusia 78 Dyskinesia(s) drug-induced 66 orofacial 66.67 tardive 66 Dysmetria 276, 277 ocular 276 Dysosmia 76 Dysphagia 102, 166, 370, 397 causes 362 Dysphonia 130 Dysplasia 288 Dysraphism, spinal 292, 293, 381 Dyssomnia 114, 116 Dyssynergy 276 Dystonia 64-65, 383 action 6 acute dystonic reactions 66. 204.205 arm 64 cervical 64.65 classification 64 craniocervical 64, 65 dopa-responsive 64 idiopathic torsion 64 leg 64 multifocal 65 oromandibular 64, 362 Parkinson disease 208, 209 paroxysmal, autosomal dominant 64 spastic 64 task-specific 64 Dystrophinopathy 336, 398 Dystrophy limb-girdle 336, 337, 400 myotonic 52, 338, 339, 400, 401 reflex sympathetic 110 see also Muscular dystrophies

E

Ecchymosis, retroauricular 267 ECG abnormalities, neurogenic 148 Echolalia 124 Edema cerebral 162, 163, 224

cytotoxic 162 hydrocephalic 162-163 treatment 264 vasogenic 162 leg. Parkinson disease and 208 Edinger-Westphal nucleus 26. 90 Edrophonium chloride test 404 Eiaculatory dysfunction 156 Elastance 162 Electro-oculography 352 Electroencephalography 352 Electrogustometry 78 Electrolyte balance disorders 310. 311 Electromyography (EMG) 352 myasthenia gravis 404 needle 391, 399 stimulation 399 urodynamic 218 Electroneurography 352 Embolism 172, 173 infectious 226 paradoxical 262 Emery-Dreifuss muscular dystrophy 336, 337 Empyema, subdural 222 **Encephalitis 222** brain stem 222 clinical manifestations 222 hemorrhagic necrotizing 236 Lyme disease and 228 toxoplasmosis and 250 viral 234, 376 herpes simplex 236 Encephalocele 292 Encephalomyelitis 222 acute demyelinating (ADEM) 234 Lyme 228 Encephalopathy 306-315 burns and 312 chronic 308 dialysis 310, 311 endocrine 310, 311 hepatic 308, 309, 387 HIV 240 hypoxic-ischemic 308, 309 iatrogenic 314, 315, 389 Lyme disease and 228 metabolic 386-387 acquired 308-312 hereditary 306-307 infancy 387 neonatal 386

mitochondrial 281 multiple organ failure and 312 paraneoplastic 312 portosystemic 308, 387 progressive myoclonic 68 septic 226, 227, 312, 313 diagnosis 226 spongiform 252-253 bovine (BSE) 252 Creutzfeldt-Jakob disease (CJD) 252, 253 genetic 252 infectious 252 subcortical arteriosclerotic (SAE) 298 substance abuse and 312. 313 trauma and 379 uremic 310, 311 Wernicke 312, 313 Endarterectomy 174 Endocarditis, bacterial 226 Endocrine encephalopathy 310, 311 myopathy 347, 398 Endolymph 56 Endoneurium 2 Endophthalmitis, candida 248 Entacapone 212 Enteroviruses 376 Enuresis 114 Eosinophilic fasciitis 344 Ependyma 6 spinal 256 Ependymoma 256, 257, 377 anaplastic 260, 377 Epidural abscess 222 hematoma 268, 270 space 6, 30 Epilepsy 192-199 acquired 198 acute epileptic reactions 196 age of onset 197 causes 198. 379 classification 196-197 epileptic syndromes 196 generalized 194, 196 genetic predisposition 198 location-related 196 mvoclonus progressive, with Lafora bodies 307 with ragged red fibers (MERRF) 403 pathophysiology 198, 199

Epilepsy prognosis 198, 373 seizure types 192-195, 373 generalized 193, 194-195, 197 grand mal 194, 195, 199 partial (focal) 192-194, 197 status epilepticus 196 grand mal 196 treatment 198 antiepileptic drugs 198 unclassified 196 see also Seizures Epineurium 2, 30 Episodic ataxia (EA) 280 headache cluster 186, 190 tension 182, 190 memory 134 paralysis 338, 339, 401 vertigo 58 Epithelium, olfactory 76 Erb palsy 318 Erectile dysfunction 156 Erythema chronicum migrans 228, 229 Esophagus 154 Essential tremor 62, 357 Estradiol 367 Ethosuximide 198 Evoked potentials 352 auditory (AEP) 218, 352 motor (MEP) 218, 352 somatosensory (SEP) 218, 352 visual (VEP) 218, 219, 352 Examination 350-351 see also specific conditions Excitotoxicity 300 **Executive functions 122** Exons 288 Expiration 150 Exteroceptors 104 Extinction phenomenon 132 Eye movements 84, 85 convergence 90 intracranial hypertension and 158 reflex 84 vergence movements 84 voluntary 84

F,

Fabry disease 307, 332 Factitious symptoms 138, 139 Falling 374 Falx cerebelli 6 cerebri 6 Famcyclovir 238 Familial spastic paraplegia (FSP) 286, 287, 384 Fascicles 2 Fasciculations 50 Fasciculus arcuate 124 cuneatus (lateral) 104 gracilis (medial) 104 longitudinal, medial 56, 84 Fasciitis, eosinophilic 344 Fatal familial insomnia 114, 280 Fatigue, multiple sclerosis 214 Felbamate 198 Festination 206 Fetal alcohol syndrome 314 Fever 152, 268 Fiber(s) commissural 24 corticopontine 44 parasympathetic 90, 140, 154, 156 preganglionic 140 sudoriparous 152 sympathetic 90, 140, 154, 156 taste 96 "U fibers" 244 Fibromyalgia 52 Fibrosarcoma 260 Fila olfactoria 76 Filum terminale 2 externum 30 internum 30 Finger agnosia 132 Finger-finger test 276, 277 Finger-nose test 276 First aid 270, 271 Fistula, dural 178, 282, 283 Fits see Seizures Fluconazole 248 Flucytosine 248 Fluid balance 143, 268, 367 disorders 310 Flunarizine 204 Fluphenazine 204 Folic acid deficiency 286 Foramen(ina)

interventricular, of Monroe 8 258 of Luschka 8 of Magendie 8 vertebral 30 Forgetfulness 134 benign senescent 134, 136 see also Memory Fornix 144, 145 Foscarnet 244 Fossa, cranial 4 anterior 4 middle 4 posterior 4, 18 Fovea 80 Fracture skull 266 vertebral 272, 380 atlantoaxial dislocation 380 bilateral axis arch 380 burst 380 compression 380 dens 380 dislocation 380 lefferson's 380 stability 272, 273 Frev syndrome 362 Friedreich ataxia 280, 281 Fronto-orbital lesions 122 Fungal infections 180, 248-249 aspergillosis 248, 249 candidosis 248, 249 cryptococcosis 248, 249 mucormycosis 248, 249

G

Gabapentin 198 Gag reflex 26 Gait 60 antalgic 60 apraxia 128, 374 ataxia 60. 61. 276-277 cycle 60, 61 disturbances 49, 60-61 factitious 139 intracranial hypertension 158 neurosyphilis 230 normal-pressure hydrocephalus 160, 161 Parkinson disease 206, 207 dystonic 60, 61 spastic 60, 61 steppage 60, 61 waddling 60

422

Galactosemia 386 Galea aponeurotica 4 Gamma-aminobutyric acid (GABA) 140, 300 Gancyclovir 244 Gangliocytoma 377 Ganglioglioma 377 anaplastic 377 Ganglioma, parasympathetic 258 Ganglion(a) basal 24, 42, 210, 211 connections 210, 211 hemorrhage 176 Parkinson disease and 210. 211 dorsal root 2 spinal 2 trigeminal central connections 94, 95 peripheral connections 94. 95 varicella-zoster and 238 Ganglioneuropathy 390 Ganglionitis 238 Gangliosidosis GM1 307, 387 GM2 307 Ganser syndrome 138 Gastrointestinal function 154-155 neurological causes of dysfunction 370 Gastroparesis 370 Gaucher disease 306, 307, 387 Gaze deviation contralateral 86 skew 70 89 Gaze palsy 88 contralateral 86 insilateral 86 Parkinson disease 208 progressive supranuclear palsy 303 Genetic predisposition epilepsy 198 Parkinson disease 213 spongiform encephalopathies 252 Genotype 288, 289 Germinoma 258, 377 Gerstmann syndrome 128 Gerstmann-Sträussler-Scheinker syndrome 280 Giant axon neuropathy 397 Gibberish, fluent 124

Gilles de la Tourette syndrome 68 Glasgow coma scale 378 Glatiramer acetate 220 Glioblastoma 260, 261, 265, 377 Glioma butterfly 260 mixed 377 Gliomatosis cerebri 260 Globus hystericus 102 pallidus 210 Glomeruli, olfactory 76 Glomus carotid 150 tumor 377 Glutamate 140, 210 antagonists 212 Glutamate decarboxylase 300 Glycerol 264 Glycogen storage disease 340 Goiter 311 Gonadotropins 143 Gordon reflex 40 Granulomatosis lymphomatoid 180 Wegener 180 Growth hormones 143, 367 GH-secreting tumors 258 Guillain-Barré syndrome 244, 326-327, 395 clinical spectrum 395 diagnosis 326, 396 pathogenesis 326 symptoms and signs 326 treatment 326 Gustatory pathway 78 Gyrus(i) angular 124 cingulate 144 dentate 144 postcentral 12 precentral 12

н

Haemophilus influenzae 376 chemoprophylaxis 226 vaccination 226 Hair cells 100 Hallucinations olfactory 76 Parkinson disease 208 Haloperidol 204 Hand grip test 369 Hangover 188 Hartnup disease 306 Head injury 267 assessment criteria 379 classification 267 complications 269, 379 see also Trauma Head, zones of 110, 111 Headache 182-191, 373 brain tumors and 254, 255 cervical syndrome 188, 189 upper 188 chronic daily 182 causes 373 chronic paroxysmal hemicrania (CPH) 186, 190 cluster 186, 187, 190 chronic 186, 190 episodic 186, 190 pathogenesis 186 combination 182 diagnostic classification 191 ice-pick 182 intracranial hypertension and 158 intracranial hypotension and 160 migraine 184-185, 190, 373 symptoms and signs 184 aura 184, 185 headache phase 184 prodromal phase 184 resolution phase 184 nociceptive transmission 188. 189 posttraumatic 270, 271 prophylaxis 190 sinus 186, 187 substance-induced 188, 189 acute 188 rebound 188 tension 182, 183 chronic 182, 190, 373 episodic 182, 190 pathogenesis 182 treatment 182, 190 trigeminal neuralgia 186. 187.190 idiopathic 186 pathogenesis 186 symptomatic 186 vascular processes and 166, 182.183 subarachnoid hemorrhage 176 Hearing 100-101 age-related changes 382 sound perception 100

Heart 148-149 Heel-knee-shin test 276 Heerfordt syndrome 362 Helicotrema 100 Hemangioblastoma 256, 257 Hemangioma 294 Hemangiopericytoma 377 Hematoma 268 epidural 268, 270 intracerebral 270 intraparenchymal 268 posttraumatic 267 subarachnoid 268 subdural 268, 270 chronic 379 Hemianopsia 82 heteronymous 82 homonymous, altitudinal 82 Hemiballism 66-67 Hemicrania, chronic paroxysmal (CPH) 186, 190 Hemineglect 132 Hemiparesis 47, 122 contralateral 46, 158 Hemiplegia 122 alternans 46 Hemodynamic abnormalities 148. 174 Hemorrhage 166, 176-179 intracerebral 166, 167, 176 basal ganglia 176 brain stem 176 caudate 176 cerebellar 176 complications 176 lobar 176 massive hypertensive 178 pathogenesis 178 pontine 176 putaminal 176 thalamic 176 treatment 178 intraventricular 166, 176, 177.270 complications 176 symptoms and signs 176 spinal 282 subarachnoid 166, 176-177, 270 complications 176 pathogenesis 178 symptoms and signs 176 treatment 178 see also Stroke Hepatic encephalopathy 308, 309, 387 Hereditary diseases 288

diagnosis 288 inheritance 288 289 mutations 288 phenotype 288 see also specific diseases Hereditary hemorrhagic telangiectasia (HHT) 294 Hereditary motor-sensory neuropathy (HMSN) 332, 333.396 type I 332, 333, 396 type II 332, 396 type III 332, 396 Hereditary neuropathy with pressure palsies (HNPP) 332. 333 Hering-Breuer reflex 150 Herniation intervertebral disk 318, 319, 392 intracranial hypertension and 158, 159, 162 subfalcine 162 transtentorial 118, 158 Herpes labialis 236 Herpes simplex 236-237, 376 pathogenesis 236 symptoms and signs 236 treatment 236 type 1 (HSV-1) 236, 237 type 2 (HSV-2) 236 Herpes zoster 180, 238, 239 occipitocollaris 238 ophthalmicus 238 oticus 238 sine herpete 238 symptoms and signs 238 Hiccups 68 Hippocampus 144, 145 Histiocytoma, malignant fibrous 260 History taking 350 Holmes tremor 357 Homocystinuria 307 Hormones aglandotropic 142 glandotropic 142 growth 143 thyroid 143 see also specific hormones Horner syndrome 48, 92, 152, 318 central 92 Hospital hopper 138 Human immunodeficiency virus (HIV) 240-241, 376 antiretroviral therapy 240

pathogenesis 240, 241 primary infection 240 secondary complications 240 symptoms and signs 240 Huntington 300 Huntington disease 66, 300. 301 inheritance 300 anticipation 300 pathogenesis 300 symptoms and signs 300 Westphal variant 300 Hydranencephaly 290, 381 Hvdrocephalus 161-163, 290, 291, 366, 379, 381 acute 162, 290 brain edema 162-163 chronic 162 communicating/malresorptive 162 congenital 290 external 162 non-communicating/obstructive 162, 258, 290 normal-pressure 160, 161, 162 subarachnoid hemorrhage and 176 symptoms and signs 290 treatment 290 Hydrochlorothiazide 338 Hydrophobia 246, 247 Hvdroxocobalamin 286 Hygroma, subdural 379 Hyperalgesia 316 Hyperammonemia 386 Hypercalcemia 310 Hypercapnia 308 Hypereosinophilia syndrome 344 Hyperesthesia 316 Hyperglycemia 308, 324 hyperosmolar nonketonic 308 Hyperglycinemia, nonketonic 386 Hyperhidrosis, Parkinson disease and 208 Hyperkalemia 52, 338, 401 Hyperkinesia 383 Hypermetria 70 Hyperprolactinemia 258 Hypersomnia 114, 116 Hypertension 148 intracerebral hemorrhage and 178

intracranial 158-160, 224, 268 290 causes 371 treatment 264, 270 Hyperthermia central 152 malignant 346-347 Hypertrophy, muscular 334, 338, 397 Hyperventilation syndrome 204, 205, 370 Hypervolemia 310 Hypesthesia 106 Hypocalcemia 310 Hypochondriacal disorder 138. 139 Hypogeusia 78 Hypoglycemia 308, 309, 324 acute 308 chronic 308 subacute 308 Hypokalemia 52, 338 Hypokinesia 206 Hypomagnesemia 310 Hypometria 70 Hypomimia 206, 207, 362 Hypoperfusion 174 Hypophonia 206 Hyposmia 76 Hypotension 148, 268 antiparkinsonian medications and 208 idiopathic orthostatic 302 intracranial 160-161 causes 372 Hypothalamic-pituitary regulatory axis 367 Hypothalamus 24, 140, 142-143 functions 142 fluid balance 310 thermoregulation 152 neuroendocrine control 142, 143 pain and 108 Hypothermia 152 Hypothyroidism 311 Hypoventilation 370 Hypovolemia 310 Hypoxia 268 global cerebral 172 hypoxic-ischemic encephalopathy 308, 309

1

Iatrogenic encephalopathy 314. 315, 389 Idiopathic cerebellar ataxia (IDCA) 276 orthostatic hypotension 302 Parkinson disease 302 torsion dystonia 64 trigeminal neuralgia 186 Immunization see Vaccination Incisura, tentorial 6 Inclusion body myositis 52. 252, 344, 345 Incontinence fecal 216, 370 urinary see Bladder dysfunction Incus 100 Infantile cerebral palsy see Cerebral palsy Infarction 172, 174 anterior cerebral artery 168 anterior choroidal artery 168 border zone 168, 172, 173 cerebellar arteries 170 dementia and 298 multi-infarct dementia 298 strategic infarct dementia 298 dorsolateral 170 end zone 172, 173 hemodynamic 172 internal carotid artery 168, 169 border zone 168 territorial 168 lacunar 168, 172 low-flow 172 paramedian 170 pontine 170 spinal central 282 complete 282 territorial 168, 172, 173, 224 hemorrhagic conversion 172 threshold 174 types of 172 Infections see Central nervous system (CNS); specific infections Inflammatory response 224 multiple sclerosis 220 tuberculous meningitis 232 Infundibulum 6

Inheritance 288, 289 monogenic 288 multifactorial 288 polygenic 288 Innervation ratio 44 Insomnia 114 fatal familial 114, 280 psychogenic 114 Inspiration 150 Interoceptors 104 Intervertebral disks 30 Intestine colon 154 pseudo-obstruction 370 small 154 Intoxication 312, 313 Intracranial pressure (ICP) 158-163 brain tumors and 254 compliance 162 elastance 162 hypertension 158-160, 268, 200 causes 371 clinical features 158-160 treatment 264, 270 hypotension 160-161 causes 372 symptoms 160 pathogenesis 162 Introns 288 Involution 296 Iodine-starch (Minor) test 152 Iris 90 Ischemia 166, 268 cerebral 148 delaved 176 global 172 hypoxic-ischemic encephalopathy 308, 309 threshold 174 transient ischemic attack (TIA) 166 see also Stroke Ischemic penumbra 174 Isocortex 24 Isoniazid 232 Itraconazole 248

J

Jaw jerk reflex 26 JC virus 244 Jefferson's fracture 380 Jet lag 114 Joints, Charcot 230 Jugum sphenoidale 4

K

Karnofsky scale 26, 378 Kearns–Sayre syndrome (KSS) 403 Keratoconjunctivitis 236 Kernig's sign 222 King–Denborough syndrome 347 Klippel–Feil syndrome 292, 293 Klumpke–Dejerine palsy 318 Korsakoff syndrome 76, 308, 312 Krabbe disease 307, 332, 387 Kufs disease 307 Kussmaul's respiration 118

L

Labyrinth 56 Lacrimation, test of 98 Lacunar state 172, 173 Lacunes 172, 298 Lambert-Eaton myasthenic syndrome (LEMS) 50, 52, 342, 343, 406 Lamotrigine 198 Lance-Adams syndrome 308 Language 124-125 aphasia 124, 126-127 model 124 Larynx, voice production 130 Lasègue's sign 318 Laterocollis 64 Lathyrism 286, 304, 384 Law of Bell and Magendie 30 Leigh disease 306 maternally-inherited (MILS) 403 Lemniscus, trigeminal 94 Lennox-Gastaut syndrome 196. 198 Lens accommodation 90 Leprosy 328, 329 Leptomeninges 6 metastases 262, 263 treatment 265 Leukoaraiosis 298 Leukodystrophy globoid cell 387 metachromatic 306, 307, 332.333 Leukoencephalitis 234 Levetiracetam 198 Levodopa 208, 212 adverse effects 389

Levomepromazine 264 Lewy bodies 208, 210, 302 Lhermitte's sign 48, 49, 214 Life expectancy 296 active 296 Lifespan 296 Ligament, denticulate 30 Light reflex 90, 91 Light-near dissociation 92 Limb girdle dystrophy 336, 337, 400 Limbic system 80, 135, 144-145 functions 144 nerve pathways 144 pain and 108 structure 144, 145 syndromes 368 Lipid metabolism disorders 332.333 Lissencephaly 381 Listeria 376 Lisuride 212 Lohe frontal hemorrhage 176 lesions 122 left 122 right 122 occipital, hemorrhage 176 parietal, hemorrhage 176 temporal, hemorrhage 176 Locked-in syndrome 120, 121, 170.359 LSD abuse 314 Lumbar puncture 2, 407 intracranial hypotension and 160 multiple sclerosis and 218. 219 Lungs 150 Lyme disease 228-229 chronic 228 clinical manifestations 228. 229 diagnosis 228 pathogenesis 228 treatment 228 Lymphadenosis benigna cutis 228 Lymphoma 180, 377, 385 cerebral, primary 260, 261, 265 ocular manifestations 260 non-Hodgkin 260, 261

M

McLeod syndrome 300 Macroadenoma 258 Macrocephaly 290, 381 Macrographia 276 Macrophages 220 Maculae 56 saccular 56 utricular 56 Magnesium balance 310 Magnetic resonance imaging (MRI) 354 brain tumors 260, 264-265 multiple sclerosis 218, 219 Major histocompatibility complex (MHC) 220 Malaise, brain tumors and 254 Malaria, cerebral 250, 251 Malformations 288, 381 Chiari 292, 293 Dandy-Walker 292 Malignancy see Brain tumors; specific tumors Malignant hyperthermia 346-347 Malignant neuroleptic syndrome 208, 347 Malingering 138 Malleus 100 Mannitol 264 Maple syrup urine disease 386 Marcus-Gunn pupils 214 MDMA abuse 314 Mean arterial pressure (MAP) 162, 174 Mechanoreceptors 104, 150 Medulla, adrenal 140 Medulla oblongata 2, 26 caudal 148 lesions 70 rostral 148 syndromes 70, 73, 361 dorsolateral 361 Medulloblastoma 260, 261, 265 Megalencephaly 381 Meige syndrome 64, 362 Melkersson-Rosenthal syndrome 362 Melperone 264 Membrane(s) arachnoid 6 spinal 30 basilar 100 tympanic 100 Memory 134-135

declarative (explicit) 134, 135 disorders 134 368 Alzheimer disease 297 head trauma and 269 Parkinson disease 208 episodic 134 examination 134, 353 long-term 134 nondeclarative (implicit) 134.135 semantic 134 short-term 134 Ménière's disease 58 Meningeosis, neoplastic 262 Meninges 6-7 Meningioma 256, 257, 377 anaplastic 377 extracranial 256 familial 256 treatment 264 Meningism 222 Meningitis 222, 268 aseptic 234, 236, 242 postinfectious 234 postvaccinal 234 asymptomatic 230 bacterial 180, 226, 376 borrelia-related (Lyme) 228 tuberculous 232-233 pathogenesis 232 symptoms and signs 232.233 treatment 232 Candida and 248 carcinomatous 262 chronic 232 clinical manifestations 222 Lyme disease and 228 Mollaret 234 neurosyphilis and 230, 231 prophylaxis 226 viral 234, 376 pathogens 234 poliovirus 242 Meningococcus 376 vaccination 226 Meningoencephalitis 222 arteritis and 226 bacterial 226, 376 Candida and 248 cryptococcosis and 248 neurosyphilis and 230 prophylaxis 226 treatment 224 guidelines 375 tuberculous 232

viral 234-235 herpes zoster 238 pathogens 234 Meningopolyradiculitis 230 Mental retardation 381 Mesencephalon 2, 26 Metachromatic leukodystrophy 306, 307, 332, 333 Metastatic disease 262-263 cascade hypothesis 262 drop metastasis 260, 262 intracranial 262, 263, 265 spinal 262, 263, 265 treatment 265 Methotrexate 220 Methylprednisolone 220, 238 Metoclopramide 204 Mexiletine 338 Microadenoma 258 Microaneurvsm 179 Microangiopathy 172 Microcephaly 381 Microglia 220 Micrographia 128, 206, 207 Micturition 156 Migraine 184-185, 373 pathogenesis 184 symptoms and signs 184 aura 184, 185 headache phase 184 prodromal phase 184 resolution phase 184 treatment 190 Migration disorder 381 Miller-Fisher syndrome 327, 395 Mini-Mental Status Examination 136 Mini-syndrome test 136 Minor test 152 Miosis 90, 382 unilateral 92 Mitochondrial disorders 288, 307.398 encephalopathy 281 myopathy 52, 398, 403 Mitoxantrone 220 Möbius syndrome 362 Mollaret meningitis 234 Monoclonal gammopathy of undetermined significance (MGUS) 328 Mononeuritis multiplex 240 Mononeuropathy 50, 180, 316, 318, 322-323, 390, 394 multiplex 316, 390 see also Neuropathy

Monoparesis 46, 47 Monoradiculopathy 316, 318 lumbar 318 Monto-Kelli doctrine 162 Motor end plate 2 lesions 50 Motor evoked potentials (MEP) 218, 352 Motor function age-related changes 382 Parkinson disease and 210 Motor neuron diseases 304-305.384-386 lower 50, 304, 385, 386 treatment 304 upper 46, 304, 384, 386 Motor unit 44 Movements mass 46 periodic leg 114 reflex 42 respiratory 150, 151 rhvthmic 42 spontaneous 50 coma staging 118 voluntary 42 see also Dyskinesia(s): Dystonia; Eye movements; Tics; Tremor Mucor (mucormycosis) 248 rhinocerebral 248, 249 Multi-infarct dementia 298 Multifocal motor neuropathy (MMN) 328, 329 Multiple organ failure 312 Multiple sclerosis (MS) 214-221 clinical manifestations 214-217. 218. 219 autonomic dysfunction 216, 217, 371 behavioral changes 216, 217 fatigue 214 incoordination 216, 217 pain 214 paresis 214 paroxysmal phenomena 216.217 sensory disturbances 214, 215 spasticity 214 visual impairment 214, 215 course of 214, 216 benign 216 chronic progressive 214 malignant 216 relapsing-remitting 214 diagnostic criteria 375

Index

Multiple sclerosis (MS) differential diagnosis 216 laboratory tests 218 bladder function 218 cerebrospinal fluid examination 218, 219 evoked potentials 218 neuroimaging 218, 219 pathogenesis 218-221 activation 220, 221 antigen presentation and stimulation 220 demvelination 220 passage through bloodbrain barrier 220 scar formation 220 prognosis 216 rehabilitation 220 relapse 214, 220 remission 214 treatment 220 symptomatic therapy 220 Multiple system atrophy (MSA) 276, 302, 303 bladder dysfunction and 371 Mumps virus 376 Münchhausen syndrome 138 by proxy 138 Muscle phosphorylase deficiency 402 Muscle(s) atrophy 49-52, 107, 281, 287, 334, 406 peripheral neuropathy and 316 poliomvelitis and 242. 243 spinal (SMA) 385 spinobulbar 385 cramps 397 detrusor 156 expiratory, auxiliary 150, 151 facial, syndromes affecting 362 functional disorders 52 hypertrophy 334, 338, 397 levator palpebrae superioris 90 pain 52 see also Myalgia respiratory 150 auxiliary 150 segment-indicating 32, 357 spasms 334 stiffness 52, 334 tarsal, superior 90

tone 276 weakness 334, 374, 397, 405 see also Myopathy Muscular dystrophies 336-337, 398.400 Becker 336, 337, 400 diagnosis 336 Duchenne 52, 53, 336, 337, 400 Emerv-Dreifuss 336, 337 facioscapulohumeral 337, 400 pathogenesis 336, 337 symptoms and signs 336 treatment 336 see also Dystrophy Mutation 288 chromosome 288 gene 288 genome 288 germ-line 288 somatic 288 Myalgia 52, 334, 346, 397 causes 346 toxic 405 Myasthenia gravis 52, 342-343, 406 aggravating drugs 403 crises 404 diagnosis 342 ancillary tests 404 pathogenesis 342 symptoms and signs 342 treatment 342 Mycobacterium leprae 328 tuberculosis 232, 376 Mycosis, opportunistic systemic 248 see also Fungal infections Mvdriasis 90 unilateral 92 Myelin sheath 2 lesions, multiple sclerosis 220 Myelinolysis, central pontine 310. 315 alcoholism and 314 Myelinopathy 316 Myelitis 222, 282 clinical manifestations 222 Lyme disease and 228 treatment 286 viral 282, 376 herpes simplex 236 herpes zoster 238 Myelography 354

Myelomeningoradiculitis, tuberculous 232 Myelopathy 282–287 acute 282-283 cervical 284, 285 chronic 48, 284-284 diagnostic studies 286 hereditary 186 HIV 240 subacute 284-285 subacute combined degeneration (SCD) 286, 287 toxic 286. 287 treatment 286 Mvoclonus 66-69, 383 epilepsy with ragged red fibers (MERRF) 403 essential 68 myoclonic encephalopathies 68 physiological 68 progressive myoclonus epilepsy with Lafora hodies 307 sleep 68, 114 symptomatic 68 Myokymia 50 Myopathy 50-52, 334-347, 397-405 carnitine deficiency 304, 402 causes 334 centronuclear 402 congenital 52, 340, 341, 398, 402 critical illness (CIM) 347, 379 diagnosis 334, 399 endocrine 347, 398 episodic paralyses 338, 339 hereditary 398 inflammatory 344, 398 malignant hyperthermia 346-347 metabolic 340, 341, 402 mitochondrial 52, 340, 341, 398, 403 mvalgia 346 myasthenic syndromes 342-343 myotonias 338, 339, 401 nemaline 402 paraneoplastic syndromes 347 primary 52 rhabdomyolysis 346 secondary 52 symptoms and signs 334 toxic 398, 405

neuromuscular syndromes 347 398 see also Muscular dystrophies Myopathy, encephalopathy, lactic acidosis and strokelike episodes (MELAS) 403 Myositis 52, 344-345 diagnosis 344 inclusion body 52, 252, 344, 345 infectious 344 Lyme disease and 228 ossificans 379 pathogenesis 344 syndromes 344 toxoplasmosis and 250 treatment 344 viral 52 Mvotomes 32, 33 Myotonia 338, 339, 401, 405 action 338 congenital 52, 338, 339 Becker 401 Thomsen 401 fluctuans 401 paradoxical 338 pathogenesis 338 percussion 338 symptoms and signs 338 treatment 338 Myotonic dystrophy 52, 338, 339, 400, 401

Ν

Nacrolepsy 114 Nasal cavity 4 Natalizumah 220 Nausea brain tumors and 254, 255 intracranial hypertension and 158, 159 Near response 90 Neck stiffness, meningitis and 222, 223 Necrosis 174 focal 224 Needle electromyography 391, 399 Neocerebellum 54 Neologisms 124, 126 Nerve injuries 330, 331 pathogenesis 330 compression 330 crushing injuries 330 transection 330

Nerve palsy abducens 86 87 facial 96, 98, 99 Lyme disease 228, 229 oculomotor 86, 87, 92, 158 trochlear 86, 87 complete 86, 87 incomplete 86 Nerve roots dorsal 2 spinal 30 root filaments 30 trauma 272 avulsion 330 ventral 2 spinal 30 Nerve(s) abducens, palsy 86, 87 alveolar inferior 94 superior 94 auriculotemporal 94 axillary 35 mononeuropathy 322, 394 buccal 94 cochlear 100 cranial 2, 6, 28-29, 356 herpes zoster and 238 motor cranial nerve nuclei 130 nerve pathways 26, 28 see also specific nerves cutaneous, lateral, of the thigh 37 mononeuropathy 323, 394 ethmoid anterior 94 posterior 94 facial 96-99 functional systems 96 lesions 98-99 examination 98 palsy 96, 98, 99 nerve pathways 96 femoral 37 mononeuropathy 323, 394 frontal 94 gluteal, mononeuropathy 323 infraorbital 94 lacrimal 94 lingual 94 mandibular 94 maxillarv 94 median 35 mononeuropathy 322, 394 meningeal 94

mental 94 musculocutaneous 35 nasociliary 94 obturator, mononeuropathy 323 oculomotor, palsy 86, 87, 92, 158.325 olfactory 2 ophthalmic 94 optic 2, 372 atrophy 158, 159 peripheral 2 peroneal 37 mononeuropathy 323, 394 phrenic 150 radial 35 mononeuropathy 322, 394 sciatic 37 mononeuropathy 394 spinal 2, 31, 32, 150 supraorbital 94 supratrochlear 94 sural, biopsy 391 thoracic, long, mononeuropathy 322, 394 tibial 37 mononeuropathy 323, 394 trigeminal 6, 76, 94-95 dysfunction 98 mesencephalic nucleus 94 motor nucleus 94 principal sensory nucleus 94 spinal nucleus 94 trochlear, palsy 86, 87 ulnar 35 mononeuropathy 322, 394 vagus 6, 154 zvgomatic 94 Neural tube defects 292, 293 Neuralgia 363 postherpetic 238 trigeminal 186, 187, 190 idiopathic 186 multiple sclerosis 214 pathogenesis 186 symptomatic 186 Neuralgic amyotrophy 321, 328. 329 Neurapraxia 330, 331 Neurites, Lewy 210 Neuritic plaques (NPs) 297 Neuritis optic, multiple sclerosis 214, 218 retrobulbar 214 toxoplasmosis and 250

Neuroacanthocytosis 300, 301 Neuroblastoma 377 Neuroborreliosis 228-229 chronic 228 clinical manifestations 228. 229 diagnosis 228 pathogenesis 228 treatment 228 Neurocranium 4 Neurocybernetic prosthesis (NCP) 198 Neurocysticercosis 250, 251 Neurodegenerative diseases 296 - 305aging 296 degenerative changes and 296 disease and 296 see also specific diseases Neurofibrillary tangles (NFTs) 297 Neurofibroma 294, 295 Neurofibromatosis 294, 295 symptoms and signs 294 type 1 (NF1) 294 type 2 (NF2) 258, 294 Neurography 391, 399 Neurohypophysis 142 Neuroimaging 353-354, 391, 399 brain tumors 254, 264-265 multiple sclerosis 218, 219 myasthenia gravis 404 Neuroleptics, adverse effects 389, 403 acute dystonic reaction 204 malignant neuroleptic syndrome 208 Neuroma, acoustic 258, 259, 294 Neuromodulators, autonomic nervous system 140-141 Neuromuscular junction 2 lesions 50, 347, 398 paraneoplastic syndromes 406 postpolio syndrome and 242 Neuromyotonia 52, 406 Neuronal ceroid lipofuscinosis 307 Neuronopathy 316, 390, 406 Neurons medium spiny-type 210

spinal parasympathetic 140 sympathetic 140 Neuropathy acquired 317, 390 acute motor-axonal (AMAN) 395 acute motor-sensory axonal (AMSAN) 395 amvloid 333 diagnosis 316, 391 giant axon 397 hereditary 317. 390 peripheral 316-333 diabetic 324, 325 hereditary 332-333 metabolic 332, 333 nonmetabolic 332, 333 infectious origin 328 inflammatory polyneuropathies 326-329 mononeuropathies 318, 322-323, 390, 394 multifocal motor (MMN) 328.329 nerve injuries 330-331 pathogenesis 330 plexopathy 318 radicular lesions 318, 320 uremic 324 vasculitic 328, 329 small fiber 316 symptoms and signs 316 autonomic 316 motor dysfunction 316 sensory dysfunction 316 syndromes 316-317, 390 tomaculous 332 Neuropeptides, pain reception and 108 Neuroprotective therapy 212 Neurosyphilis 230-231 clinical manifestations 230, 231 early meningitis 230, 231 progressive paralysis 230, 231 tabes dorsalis 230, 231 meningovascular 230 pathogenesis 230 treatment 230 Neurotmesis 330, 331 Neurotransmitters autonomic nervous system 140-141 pain reception and 108 Parkinson disease and 210

Neurotuberculosis 232 Niemann-Pick disease 306. 307. 387 Nightmares 114 Ninhvdrin test 152 Nociception 108 see also Pain Nodes of Ranvier 2 Nonneurological disorders 138-139 Norepinephrine 140, 382 Notch, tentorial 6 Nucleus(i) ambiguus 102, 148 caudate 210 cochlear anterior 100 posterior 100 cuneatus 104 detrusor 156 Edinger-Westphal 26, 90 gracilis 104 lentiform 210 motor cranial nerve 130 oculomotor 90 Onuf's 156 Perlia's 90 pulposus 30 red 24 respiratory group dorsal 150 ventral 150 solitary tract 148 subthalamic 24, 210 tractus solitarius 102 ventral posterolateral (VPL) 104 vestibular 26, 56 Nystagmus 70, 86, 88-89 congenital 88 end-position 88 examination 88 gaze-evoked 88, 89, 276, 277 gaze-paretic 88 jerk 88 multiple sclerosis 214, 215 optokinetic 84, 88 pathological 88 physiological 88 positional 70, 88 see-saw 70 spontaneous 88, 89 vestibular central 88, 89 peripheral 88, 89

Index

0

Obstructive sleep apnea 114 Occlusion 172 basilar artery 170, 171 brachiocephalic trunk 168 carotid artery common 168 internal 168 cerebellar arteries 170, 171 cerebral arterv middle 168, 169 posterior 170, 171 lenticulostriate artery 168 ophthalmic artery 168 subclavian artery 170 Ocular deviation 70 Oculomotor disturbances 86-88 276 examination 86 internuclear disturbances 86 peripheral disturbances 86 stroke and 166 supranuclear disturbances 86 Oculomotor function 84-85 Odynophagia 102 Olfaction 76-77 disturbances 76 tests of 76 Olfactory pathway 76 Oligoastrocytoma 256, 377 anaplastic 377 Oligodendroglioma 256, 377 anaplastic 260, 261, 377 One-and-a-half syndrome 86 Onuf's nucleus 156 Ophthalmoplegia chronic progressive external (CPEO) 403 internuclear 86, 87, 214 Ophthalmoscopy 80 Opiate abuse 314 Oppenheim reflex 40 **Opportunistic infections 240** fungal 248-249 aspergillosis 248, 249 candidosis 248, 249 cryptococcosis 248, 249 mucormycosis 248, 249 Orbit 4, 372 Organ(s) circumventricular 140 of Corti 100 subfornical 140 Organum vasculosum 140

Orientation evaluation 353 Orthostasis test 369 Osler–Weber–Rendu disease 294 Osmoregulation 310 Ossicles, auditory 100 Overlap syndrome 344 Oxcarbazepine 198 Oxidative stress 212

P

Pachygyria 381 Pachymeninges 6 Pain 108-111 classification of 363 complex regional pain syndrome (CRPS) 110 diabetic neuropathy and 324 dvsesthesia 106 facial, atypical 373 multiple sclerosis 214 myelopathies 282 nerve pathways 104, 109 neurosyphilis 230 Parkinson disease 208, 209 pathogenesis 108 persistent somatoform pain disorder 138, 139 processing 108, 109 neurotransmitter/neuropeptide role 108 pseudoradicular 32 radicular 32 reception 108 referred 108, 110, 363 headache 188, 189 sensitization central 108 peripheral 108 transmission 108 treatment 264, 324 types of 108, 109, 363 central 363 chronic 363 deafferentation 363 neuropathic 108, 363 nociceptive 108, 363 phantom limb 363 psychogenic 363 radicular 363 somatic 108. 363 visceral 108, 110 zones of Head 110, 111 see also Myalgia Paleocerebellum 54 Pallidotomy 212

Palsv cerebral, infantile 288-291, 381 causes 288 symptoms and signs 288 treatment 290 Erb 318 Klumpke-Dejerine 318 progressive supranuclear (PSP) 208, 302, 303 pseudobulbar 362 see also Nerve palsy Pancoast tumor 262 Panic disorder 202, 203 Papez circuit 144 Papilledema 158, 159, 160 brain tumors and 254, 255 chronic 158, 159 meningitis and 222 Papilloma, choroid plexus 256. 257.377 Paraganglioma 258 sympathetic 258 Paragrammatism 124, 126 Paralysis central 46-49 upper motor neuron (UMN) lesions 46 crossed 46, 47, 70 episodic 338, 339, 401 diagnosis 338 pathogenesis 338 prophylaxis 338 symptoms and signs 338 treatment 338 peripheral 50-53 poliomyelitis and 242, 243 progressive, neurosyphilis and 230, 231 spinal, familial spastic 286, 287 Paramyotonia cold-induced 52 congenita 338, 339, 401 Paraneoplastic syndromes 312, 347.388 neuromuscular 406 Paraparesis 46 tropical spastic 384 Paraphasia 126 phonemic 124 semantic 124 Paraplegia familial spastic (FSP) 286, 287.384 in flexion 214 spinal cord trauma and 274

Parapraxia 128 Paraproteinemic polyneuropathy 328, 329, 406 Parasomnias 114 Parasympathetic nervous system 90, 140, 147, 148, 154 Paresis crossed 47 insilateral 46 Lyme disease and 228, 229 multiple sclerosis 214 peripheral 47 Paresthesia 106, 316 multiple sclerosis 214 spinal artery syndrome 282 Parinaud syndrome 70, 92, 358 Parkinson disease 62. 206-213 agraphia 128 autonomic dysfunction 208, 209 bladder disorders 208. 371 blood pressure changes 208 constipation 208 hyperhidrosis 208 leg edema 208 seborrhea 208 sexual dysfunction 208 sleep disorders 208, 209 behavioral changes 206-209 anxiety 206 dementia 208 depression 206 hallucinations 208 cardinal manifestations 206, 207 akinesia 206 bradykinesia 206 hypokinesia 206 postural instability 206, 207 rigidity 206, 207 tremor 206, 207 dvstonia 208, 209 genetics of 213 idiopathic 302 pathogenesis 210-211 basal ganglia 210, 211 connections 210 211 motor function 210 neurotransmitters 210 sensory manifestations 208 dysesthesias 208 pain 208, 209 treatment 212-213 deep brain stimulation 212 neuroprotective 212

stereotactic neurosurgical procedures 212 symptomatic 212 transplant surgery 212 visual disturbances 208 Parkinsonism 374, 383 atypical 302-303 Parosmia 76 Paroxysmal depolarization shift 198 Pathway(s) auditory 100, 101 cranial nerves 26-28 deglutition 102, 103 direct 210 facial nerve 96 gustatory 78 indirect 210 limbic system 144 olfactory 76 pain 104, 109 pupillomotor 90 somatosensory 104 visual 80-81 Peduncle, cerebellar inferior 54 lesions 358 middle 54 superior 54 Pelizaeus-Merzbacher disease 387 Penicillin 230 Peregrinating patient 138 Pergolide 212 Pericranium 4, 6 Perineurium 2.30 Peripheral nervous system (PNS) 2.3 Peristalsis 154 Peritonitis 226 Perlia's nucleus 90 Perphenazine 204 Perseveration 126 Persistent somatoform pain disorder 138, 139 Persistent vegetative state 120, Personality 122 Phakomatoses 294-295, 381 Phencyclidine (PCP) abuse 314 Phenobarbital 198 Phenomenon black-curtain 168 doll's-eye 26, 302, 303 coma and 118 extinction 132 rebound 276, 277

Uhthoff's 214 Phenotype 288, 289 Phenylketonuria 306 Phenvtoin 198, 264 Pheochromocytoma 258 Phonation 130 dysphonia 130 Phosphofructokinase deficiency 402 Photoreceptors 104 Physical examination 350-351 Pia mater 6 spinal 30 Pick disease 298, 299 Pineal region tumors 258, 259 Pineoblastoma 258, 377 Pineocytoma 258, 377 "Pisa" syndrome 65, 66 Pituitary gland 6 anterior lobe 142 hypothalamic-pituitary regulatory axis 367 posterior lobe 142 Sheehan's postpartum necrosis 258 stalk 6 tumors adenoma 258, 259, 377 metastases 262 treatment 264 Plasmodium falciparum (cerebral malaria) 250, 251 Platybasia 292, 293 Plexopathy 50, 318, 321 Plexus(es) 32-36 brachial 32, 34, 318 infraclavicular region 318 Pancoast tumor 262 plexopathy 318, 321, 393 supraclavicular region 318 trauma 272, 330 cervical 32, 35 cervicobrachial 34 choroid 6.8 carcinoma 377 papilloma 256, 257, 377 coccygeal 32 ganglionic, submucous 154 lumbar 32, 36 lumbosacral 36 plexopathy 318, 321, 393 mventeric 154 pterygoid 20 sacral 32 lesions 318 vertebral venous external 22 internal 22

Pneumococcus 376 vaccination 226 POEMS syndrome 328 Poliomvelitis 242-243 bulbar 242 encephalitic form 242 maior 242 paralytic 242 preparalytic 242 minor (abortive) 242 pathogenesis 242, 243 postpolio syndrome 242, 243.385 prevention 242 spinal form 242 symptoms and signs 242 Poliovirus 242 Polvarteritis nodosa 180 Polymyalgia rheumatica 52 Polymyositis 52, 240, 344, 345, 405.406 Polyneuropathy 50, 316 critical illness (CIP) 347, 379 diabetic (DPN) 324, 325, 395 diagnosis 324 symptoms and signs 324 treatment 324 distal symmetric 324, 325 focal 316 hereditary, genetic features 396 hypoglycemic 395 inflammatory 326-329 paraproteinemic 328, 329, 406 sensorimotor 406 small-fiber 395 see also Neuropathy Polyradiculitis 236 Polyradiculoneuropathy 228, 406 acute inflammatory demyelinating (AIDP) 395 chronic inflammatory demyelinating (CIDP) 327, 328 HIV and 240 Polyradiculopathy 316 lumbosacral 244, 395 Pons 2, 6 hemorrhage 176 infarction 170 lesions 70 syndromes 70-72, 359-360 Pontocerebellum 54, 55 Porencephaly 290, 291, 381 Porphyria 332, 333 Postpolio syndrome 242, 243, 385

Posttraumatic syndrome 379 Posture 60 61 disturbances 106, 276 Parkinson disease and 206. 207 test 276, 277 Pramipexol 212 Praziguantel 250 Predelirium 312 Presbycusis 382 Presbyopia 382 Presenilin genes 297-298 Primary complex 232 Primary lateral sclerosis 384 Primidone 198 Primitive neuroectodermal tumor (PNET) 260, 261, 265, 377 Prion protein (PrP) 252, 253 PRNP gene 252 Progesterone 367 Progressive multifocal leukoencephalopathy (PMS) 244-245 diagnosis 244 pathogenesis 244 symptoms and signs 244 Progressive supranuclear palsy (PSP) 208, 302, 303 Prolactin 367 Prolactinoma 258 Proprioception evaluation 106 Prosencephalon 2, 3 Prosody 124 Protein X 252 Pseudo-lupus erythematosus 405 Pseudodementia 297 Pseudoinsomnia 114 Pseudoradicular syndromes 318, 320, 392 causes 320 Pseudoseizures 200 Pseudotumor cerebri 160 Pseudounipolar cells 2 Pterion 4 Pupillary dysfunction 92-93 examination 92 swinging flashlight test 92 parasympathetic denervation 92, 93 sympathetic denervation 92. 93 Pupillomotor function 90-91 pupilloconstriction 90 Pupil(s) 90-92 Argyll-Robertson 92, 230

Marcus-Gunn 214 pinpoint 92 tonic 92 Putamen 210 Pyramid, medullary 46 Pyrazinamide 232 Pyridostigmine bromide 342 Pyrimethamine/sulfadiazine 250

Q

Quadrantanopsia 82 Quadriparesis 46 Quadriplegia, spinal cord trauma and 274 Quantitative sudomotor axon reflex test (QSART) 152

R

Rabies 246-247 hyperexcitability stage 246. 247 paralytic stage 246 pathogenesis 246, 247 prodromal stage 246 prophylaxis 246 sylvatic 246 symptoms and signs 246 urban 246 Radiation, optic 80 Radiculitis 236, 238 Radiculopathy 316, 318, 320, 390 causes 320, 392 Radiography 354 Radiotherapy 264-265 adverse effects 389 Ramsay–Hunt syndrome 238 Raymond-Céstan syndrome 360 Reactions acute dystonic 66, 204, 205 acute epileptic 196 Reading 124, 125 alexia 128 Rebound phenomenon 276, 277 Receptor(s) 2, 104 baroreceptors 148 chemoreceptors 104, 150. 151 cutaneous 104 exteroceptors 104 interoceptors 104 mechanoreceptors 104, 150

Receptor(s) olfactory 76 photoreceptors 104 thermoreceptors 104, 152 Reflex(es) 26, 40-42 abnormalities 46, 50 acoustic 26, 96 age-related changes 382 blink 96, 98 coma staging 118 corneal 26, 96 cough 26 cutivisceral 110, 111 extrinsic 40, 42 flexor 40 Gordon 40 Hering-Breuer 150 intrinsic 40, 42 light 90, 91 masseter (jaw jerk) 26 oculocephalic 26 **Oppenheim 40** optokinetic 84 orbicularis oculi 96 orbicularis oris 96 palmomental 96 pathological 40, 46 pharyngeal (gag) 26 pupillary light 26 reflex arc 40 reflex response 40 snout 96 stapedius 98 startle 68 sucking 96 trigeminal autonomic reflex circuit 184 vasodilatory axon 110, 111 vestibulo-ocular (VOR) 26, 84 coma and 118 viscerocutaneous 110, 111 visceromotor 110, 111 viscerovisceral 110 visual blink 96 Refsum disease 332 Renal failure 310 Respiration 150-151 auxiliary muscles 150, 151 Cheyne-Stokes 118 costal 150 diaphragmatic 150 disorders 150, 151, 370 stroke and 166 Kussmaul's 118 respiratory movements 150, 151

rhythm 150 Respiratory test 369 Restless legs syndrome (RLS) 114 Reticular formation 26 medullary 26 mid brain 26 pain and 108 pontine 26 paramedian 84 Retina 80 lesions 372 Retrocollis 64 Rhabdomyolysis 346, 406 Rhabdomvosarcoma 260 Rhizopus 248 Rhombencephalon 2 Rhvthm circadian 112, 113 disturbances 114 respiration 150 Rifampicin 226, 232 Rigidity 206, 207 cogwheel 206, 207 Riluzole 304 Romberg sign 276 Ropinirol 212 Ross syndrome 92

S

Saccades 84 abnormal 70, 276, 277 Saccule 56 Sacrum 30 Salivation, test of 98 Sarcoglycanopathy 398 Sarcoidosis 234 Sarcoma cerebral, primary 260 meningeal 260 Scala tympani 100 vestibuli 100 Scalp 4, 7 injuries 266 Scaphocephaly 381 Scar formation, multiple sclerosis 220 Schellong's test (orthostasis) 369 Schizencephaly 381 Schwann cells 2 Schwannoma 258, 377 Sclerosis amyotrophic lateral (ALS) 304, 386

primary lateral 384 tuberous (TSC) 294, 295 see also Multiple sclerosis Scotoma 82, 83 central 82, 215 homonymous bilateral 82 unilateral 82 iunction 82 Scrapie 252 Seborrhea, Parkinson disease and 208 Segawa syndrome 64 Segment-indicating muscles 32.357 Seizures epileptic brain injury and 268 brain tumors and 254, 264 generalized 193, 194-195. 197 absence 194, 195 atonic 194 grand mal 194, 195, 199 mvoclonic 194 tonic-clonic 194, 195 isolated nonrecurring 196 partial (focal) 192-194, 197 complex 192, 193, 194 secondary generalization 192 simple 192, 193, 194 pathophysiology 198, 199 postictal period 192 status epilepticus 196 grand mal 196 stroke and 166 tuberous sclerosis and 294 types of 192-195 nonepileptic 200-205 acute dystonic reaction 66, 204, 205 drop attacks 204, 205 hyperventilation syndrome (tetany) 204, 205 panic attacks 202, 203 pseudoseizures 200 psychogenic 200, 202, 203 postictal phase 202 premonitory signs 202 semiology 202 simulated 202 syncope 200-201 tonic spasms 204, 205 Selegiline 212 Senescence 296

Sensation 104-105 enicritic 104 protopathic 104 somatic 104 superficial 106 Sensory disturbances 106-107 age-related 382 examination 106 interpretation 106 location 106, 316, 317 peripheral neuropathies 316, 318.390 radicular 107 stroke and 166 Sensory receptors see Receptors Sepsis 312, 313, 347 candida 248 see also Bacterial infections Sexual behaviour disturbances 368 Sexual function 156 multiple sclerosis and 216 Parkinson disease and 208 Shingles see Herpes zoster Shock, spinal 48, 274 Shulman disease 344 Shy-Drager syndrome 302 Sign(s)Babinski 40, 49 Bragard's 318 Brudzinski's 222 Kernig's 222 Lasègue's 318 Lhermitte's 48, 49, 214 Romberg 276 Sinus(es) carotid 10, 148 massage 369 paranasal 4 ethmoid 4 frontal 4 maxillary 4 sphenoid 4 thrombosis 180, 181 aseptic 180, 181 etiology 180 septic 180 symptoms and signs 180 treatment 180 venous dural 6. 18 thrombophlebitis 226, 227 Sinusitis 186 Skull 4-5 base 4.5 metastases 262

syndromes 74-75 fracture 266 Sleep 112-113, 363 age changes 112, 113 circadian rhythm 112, 113 disturbances 114 disorders 114-115 neurogenic 114 Parkinson disease 208. 209 primary (dyssomnias) 114 extrinsic 114 intrinsic 114 parasomnias 114 psychogenic 114 secondary 114 systemic disease and 114 non-rapid eve movement (NREM) 112, 363 polygraphic recordings 112 profile 112, 113 rapid eye movement (REM) 112.363 Smell 76-77 age-related changes 382 olfactory disturbances 76 tests of 76 Sodium balance 310, 311 Sodium channel dysfunction 338, 398 Somatization disorder 138 Somatoform disorders 138 persistent somatoform pain disorder 138, 139 Somatosensory evoked potentials (SEP) 218, 352 Somnambulism 114 Somnolence 116, 117 Sound perception 100 Space epidural 6, 30 infratentorial 6 subarachnoid 6, 8 pathogen entry 224 spinal 30 subdural 6 supratentorial 6 Virchow-Robin 6 Space-occupying lesions, intracranial pressure and 162-163 Spasm multiple sclerosis and 214 muscle 334 right hemifacial 99 tonic 204, 205 Spasmus nutans 88

Spasticity 46, 47 multiple sclerosis 214 Spatial orientation disturbances 132-133 Spectral analysis 100, 101 Speech 130 articulation 130 disorders 124, 130 dysarthria 124, 130, 276 dysphonia 130 neural basis of 130, 131 voice production 130 Sphenoid bone 4 Spielmeyer-Vogt syndrome 307 Spina bifida 292, 293 Spinal automatisms 46 Spinal cord 2, 30-32 blood supply 22-23 claudication 282 infarction central 282 complete 282 lesions 48, 275, 381 bladder dysfunction and 371 cervical 48 lumbar 48 sacral 48 thoracic 48 transection 48, 274, 282, 381 trauma 274-275, 371 acute stage (spinal shock) 274 chronic stage-late sequelae 274 closed 274 open 274 rehabilitation stage 274 treatment 380 Spinal dysraphism 292, 293 Spinal muscular atrophy (SMA) 385 Spinal shock 48, 274 Spine 30-31 hemorrhage 282 metastases 262, 263 treatment 265 neoplasms 284, 285 trauma 272-273, 380 brachial plexus 272 diagnosis 272 nerve roots 272 treatment 380 vertebral fracture 272 whiplash injury 272, 273, 380

Spinocerebellar ataxia (SCA) 280 384 Spinocerebellum 54, 55 Spironolactone 338 Split-brain syndrome 24 Spondylolisthesis 392 Spondylosis deformans 392 Spongiform encephalopathies 252-253 hovine 252 Creutzfeldt-Jakob disease (CID) 252, 253 genetic 252 infectious 252 Stapes 100 Staphylococcus 376 Startle reflex 68 Status epilepticus 196 grand mal 196 Steele-Richardson-Olszewski syndrome 302 Stem cell transplantation, Parkinson disease 212 Stenosis craniostenosis 381 spinal 392 lumbar 284 Stomach 154 Strategic infarct dementia 298 Striatal foot 64 Striatonigral degeneration (SND) 302 Stroke 166-181 bladder dysfunction and 371 causes 167. 172 hemorrhage 166, 176-179 intracerebral 166, 176 basal ganglia 176 brain stem 176 cerebellar 176 lobar 176 pathogenesis 178 intraventricular 166, 176 complications 176 symptoms and signs 176 subarachnoid 166, 176-177 complications 176 pathogenesis 178 symptoms and signs 176 treatment 178 in evolution 166 ischemia 166-175 carotid artery territory 168 pathophysiology 174, 175

hemodynamic insufficiency 174 hypoperfusion 174 prevention primary 174 secondary 174 risk factors 172 treatment 174 treatment window 174 vertebrobasilar territory 170 - 171maior 166 minor 166 sinus thrombosis 180, 181 etiology 180 symptoms and signs 180 treatment 180 symptoms and signs 166 Stupor 116, 117 Sturge-Weber disease 294. 295 Subacute combined degeneration (SCD) 286, 287 Subcortical arteriosclerotic encephalopathy (SAE) 298 Subdural empyema 222 hematoma 268, 270 chronic 379 hygroma 379 space 6 Subependymoma 256 Substance abuse 312, 314, 315 see also Alcoholism Substantia nigra 24, 210 Sucking reflex 96 Superoxide dismutase 1 (SOD1) 304 Suture(s) 4 coronal 4 lambdoid 4 sagittal 4 Swallowing see Deglutition; Dysphagia Sweating disturbances 152 sweat gland innervation 152, 153 Swinging flashlight test 92 Sympathetic nervous system 90, 140, 147, 148, 154 Sympathetic skin response (SSR) 152 Symptoms nature of 350 onset 350 severity of 350

time course 350 see also specific conditions Synchondroses 4 Syncope 200-201 causes 374 neurocardiogenic 148 Syndrome(s) acquired immunodeficiency (AIDS) 240 Adie 92 alcohol withdrawal 312, 313 alien hand 24, 302 anterior horn 48, 50 anterior spinal artery 48 Anton 132 apallic 117, 120, 121 apraxia-like 128 Bannwarth 228 Bassen-Kornzweig 300 Benedict 70, 358 brain stem 70-71 Brown-Séguard 48 carpal tunnel 322 cauda equina 319 cavernous sinus 75 central cord 48 central salt-wasting 310 cervical 188, 189 upper 188 chiasm 75 Churg-Strauss 180, 344 clivus 75 comalike 120-121 complex regional pain syndrome (CRPS) 110 conus medullaris 48 Crow-Fukase 328 decerebration 46, 47, 118, 158 159 decortication 46, 47, 118 disconnection 24, 128 disequilibrium 310 encephalitic 222 epileptic 196 fetal alcohol 314 Frev 362 frontal lobe 122 lateralized 122 nonlateralized 122 Ganser 138 Gerstmann 128 Gerstmann-Sträussler-Scheinker 280 Gilles de la Tourette 68 Guillain-Barré 244, 326-327, 395-396 Heerfordt 362

herniation 158, 162 Horner 48, 92, 152, 318 central 92 hypereosinophilia 344 hyperventilation 204, 205 Kearns-Sayre 403 King-Denborough 347 Klippel-Feil 292, 293 Korsakoff 76, 308, 312 lacunar 359 Lambert-Eaton 50, 52, 342, 343, 406 Lance-Adams 308 Lennox-Gastaut 196, 198 limbic 368 locked-in 120, 121, 170, 359 McLeod 300 malignant neuroleptic 208, 347 medullary 70, 73, 361 dorsolateral 361 Meige 64, 362 Melkersson-Rosenthal 362 meningitic 222, 234 mid brain 70, 71, 358-359 Miller-Fisher 327, 395 Möbius 362 Münchhausen 138 mvelitic 222 myopathy 397 myositis 344 neuropathy 316-317, 390 of inappropriate ADH secretion (SIADH) 310 olfactory nerve 75 one-and-a-half 86 orbital apex 75 overlap 344 paraneoplastic 312, 347, 388 neuromuscular 406 parasagittal cortical 46 Parinaud 70, 92, 358 Parkinsonian, atypical 302-303 "Pisa" 65, 66 POEMS 328 pontine 70-72, 359-360 lateral pontomedullary 70 postanoxic 308 delayed 308 posterior column 48 posterior cord 48 posterior horn 48 postpolio 242, 243, 385 posttraumatic 379 postviral fatigue 52 pseudoradicular 318, 320, 392

radicular 50, 392 Ramsav-Hunt 238 Raymond-Céstan 360 restless legs (RLS) 114 Ross 92 Segawa 64 Shv-Drager 302 skull base 74-75 sphenoid wing 75 Spielmeyer-Vogt 307 spinal artery anterior 282 posterior 282 spinal cord transection 274 split-brain 24 Steele-Richardson-Olszewski 302 Sturge-Weber 294, 295 sulcocommisural artery 282 tethered cord 292, 293 top of the basilar 70, 359 transverse cord 180, 232, 380. 381 sensory 214 Vogt-Kovanagi-Harada 234 Wallenberg 70, 170, 361 Weber 70, 358 Wernicke-Korsakoff 312. 314 West 196 Zellweger 386 Synkinesia 46, 99, 362 Syphilis 180 latent 230 see also Neurosyphilis Svringomvelia 107, 284, 285 System craniosacral 140 dioptric 80 epicritic/lemniscal 104 limbic 80, 135, 144-145 functions 144 nerve pathways 144 pain and 108 structure 144, 145 syndromes 368 protopathic 104 reticular activating (RAS) 116 spinocerebellar 104 thoracolumbar 140, 152 trigeminovascular 184, 185

T

T lymphocytes, multiple sclerosis and 220 Tabes dorsalis 230, 231 Taenia solium (neurocysticercosis) 250, 251 Takayasu arteritis 180 Tangier disease 332 Tapeworm infection 250, 251 Taste 78-79 age-related changes 382 disturbances 78 tests of 78 Taste buds 78 Tav-Sachs disease 387 Tela choroidea 6 Telangiectasia 280, 281, 294, 295 hereditary hemorrhagic (HHT) 294 Telencephalon 2 Temperature see Thermoregulation Temporal arteritis 180 Temporomandibular joint 4 Tension headache see Headache Tentorium cerebelli 6 Testosterone 367 Test(s) Aachen aphasia 124 apnea 364 Bárány's pointing 276 caloric 26 confrontation test 82, 83 edrophonium chloride 404 finger-finger 276, 277 finger-nose 276 hand grip 369 heel-knee-shin 276 iodine-starch (Minor) test 152 lacrimation 98 neurophysiological 352 neuropsychological 352-353 ninhydrin test 152 olfaction 76 posture 276, 277 quantitative sudomotor axon reflex test (QSART) 152 red vision test 82 respiratory 369 salivation 98 Schellong's (orthostasis) 369 Schirmer test 98 smell 76 swinging flashlight test 92 sympathetic skin response (SSR) 152 taste 78 Wada 126

Tetany 204 Tethered cord syndrome 292, 293 Thalamus 24 hemorrhage 176 pain and 108 Thermoreceptors 104, 152 Thermoregulation 152-153 disturbances 152 neural control 152 Thiamin deficiency 312 Thromboembolism, infectious 226 Thrombophlebitis, bacterial 226, 227 Thrombosis 172, 173 prophylaxis 264 see also Occlusion Thyroid hormones 143 Tiagabine 198 Tic douloureux 186 Tics 68-69, 362, 383 Tobacco-alcohol amblyopia 314 Toe-walking 60 Tonotopicity 100, 101 Top of the basilar syndrome 70.359 Topiramate 198 Torcular Herophili 18 Torticollis 64 Toxoplasma gondii (toxoplasmosis) 250, 251 bradyzoites 250 congenital 250 oocvsts 250 tachyzoites 250 Tract(s) corticobulbar 44 corticonuclear 96 corticopontine 44 corticospinal anterior 44 lateral 44 motor 44 pyramidal 44, 45 reticulospinal 44 rubrospinal 44 spinocerebellar 104 spinothalamic anterior 104 lateral 104 posterior 104 trigeminocortical 94 vestibulospinal 44 Transcranial Doppler (TCD) 353

Transient ischemic attack (TIA) 166 crescendo 166 Transient monocular blindness 168 372 Transmissible spongiform encephalopathies (TSEs) 252-253 Creutzfeldt-Jakob disease (CID) 252, 253 Transverse cord syndrome 180, 232, 380, 381 sensorv 214 Trauma 266-275, 366 multiorgan 266 spinal 272-273 brachial plexus 272 diagnosis 272 nerve roots 272 treatment 380 vertebral fracture 272 whiplash injury 272, 273, 380 spinal cord 274-275 acute stage (spinal shock) 274 chronic stage-late sequelae 274 rehabilitation stage 274 treatment 380 traumatic brain injury (TBI) 266 - 271complications 269, 270 evaluation 266 pathogenesis 270, 271 primary injury 266, 270 prognosis 268 secondary sequelae 268, 270 treatment 270 hospital 270 scene of accident 270. 271 types of 266 Tremor 62-63 action 62.63 cerebellar 357 essential 62, 357 genesis of 62 Holmes 357 intention 62, 63, 276 kinetic 62.63 palatal 357 Parkinson disease and 206. 207, 357 physiological 357 polyneuropathic 357

postural 62 psychogenic 357 rest 62, 63 task-specific 62 types of 63, 357 Treponema pallidum 230 Triflupromazine 204 Triiodothyronine 367 Tropical spastic paraparesis 384 Trunk brachiocephalic 10, 148 occlusion 168 lumbar sympathetic, lesions 318 Tuberculoma 232, 233 Tuberculosis 180 organ 232 reactivated 232 spinal 232 Tuberculous meningitis 232-233 diagnosis 232 pathogenesis 232 symptoms and signs 232. 233 treatment 232 Tuberous sclerosis (TSC) 294, 295 Tumors Pancoast 262 paraneoplastic encephalopathy 312 primitive neuroectodermal (PNET) 260, 261, 265, 377 spinal 284, 285 see also Brain tumors; specific tumors Turicephaly 381

U

Uhthoff's phenomenon 214 Ulegyria 381 Ultrasonography 353 duplex sonography 353 transcranial Doppler (TCD) 353 Uremia 310, 311 Uremic encephalopathy 310, 311 neuropathy 324 Urinary tract infection, multiple sclerosis 216 Utricle 56

V

Vaccination Haemophilus influenzae 226 Lyme disease 228 meningococcus 226 Pneumococcus 226 poliomvelitis 242 rabies 246 Valacyclovir 238 Valproic acid 198 Valsalva maneuver (VM) 369 Varicella-zoster 238-239 complications 238, 239 ganglionic latency phase 238 immunocompromised patients 238 pathogenesis 238, 239 symptoms and signs 238, 239 treatment 238 viral reactivation 238 Vascular reserve 174 Vasculitic neuropathy 328, 329 Vasculitides primary 180 secondary 180 Vasculitis bacterial 180, 226 cerebral 180, 181 causes 180 differential diagnosis 216 immune 181 Lyme disease and 228 primary 180 secondary 180 symptoms and signs 180 treatment 180 CMV 244 neurosyphilis and 230 Vasospasm 176 Vein(s) azvgos 22 basal (of Rosenthal) 18 bridging 18 central 18 cerebral 18-19 deep 10, 18 great (of Galen) 18 inferior 18 internal 18 superficial 10, 18 middle 18 superior 18 thrombophlebitis 226 cervical 20 deep 20, 22 cortical 18

cranial 20 diploic 18 emissary 18 facial 20 hemiazygos 22 iliac, common 22 intercostal, posterior 22 jugular external 20 internal 20 lumbar 22 occipital 18, 20 radicular 22 retromandibular 20 sacral lateral 22 medial 22 spinal 22, 23 anterior 22 lateral 22 posterior 22 vertebral 20, 22 Vena cava, superior 22 Ventriculitis 222, 226, 227 Ventriculomegaly, long-standing overt (LOVA) 290 Vertebrae 30 cervical 30 coccygeal 30 fracture 272, 380 burst 380 compression 380 dens 380 dislocation 380 Jefferson's 380 stability 272, 273 lumbar 30 thoracic 30 Vertebral column 30 Vertigo 58-59 benign paroxysmal positional 58, 59 brain tumors and 254, 255 episodic 58 nonvestibular 58, 59 physiological 58 vestibular central 58 peripheral 58 Vestibular system 56-57 disorder 374 Vestibulocerebellum 54-56 Villi, arachnoid 8 Viral infections 234-247, 376. 385 cytomegalovirus (CMV) 244-245

herpes simplex 236-237 human immunodeficiency virus (HIV) 240-241 meningoencephalitis 234-235 pathogens 234 poliomvelitis 242-243 progressive multifocal leukoencephalopathy (PMS) 244-245 rabies 246-247 varicella-zoster 238-239 see also specific infections Virchow-Robin space 6 Viremia primary 238 secondary 238 Viscerocranium 4, 5 Vision color 80 disturbances 80 intracranial hypertension and 158 stereoscopic 80 Visual disturbances color vision 80 multiple sclerosis 214, 215 Parkinson disease 208 stroke and 166 see also Visual field Visual evoked potentials (VEP) 218, 219, 352 Visual field 80 binocular 80 confrontation test 82, 83 defects 82-83 chiasmatic lesions 82 perichiasmatic lesions 82 retrochiasmatic lesions 82 monocular 80 red vision test 82 Visual pathway 80-81 Vitamin B1 deficiency 312 Vitamin B12 deficiency 286, 287 Vitamin E deficiency 280 Vogt-Koyanagi-Harada syndrome 234 Voice production 130 timbre 130 volume 130 whisper 130 see also Speech Volume regulation 310 Vomiting 370 brain tumors and 254, 255

Vomiting

intracranial hypertension and 158 von Heubner angiitis 230 von Hippel-Lindau disease 294, 295 hemangioblastoma and 256 symptoms and signs 294 von Recklinghausen disease 294

W

Wada test 126 Wakefulness 116 *see also* Consciousness Wallenberg syndrome 70, 170, 361 Water balance see Fluid balance Watershed zones 22, 168 Weber syndrome 70, 358 Wegener granulomatosis 180 Wernicke-Korsakoff syndrome 312.314 Wernicke's aphasia 126, 127 area 124 encephalopathy 312, 313 West syndrome 196 Whiplash injury 272, 273 severity classification 272 symptoms and signs 272 treatment 380 Whisper 130

White-matter lesions (WMLs) 298 Wilson disease 208, 307 Writer's cramp 64, 65 Writing 124 agraphia 128, 129

Х

Xanthoastrocytoma, pleomorphic 256, 377

Ζ

Zellweger syndrome 386 Zones, of Head 110, 111