To Sue
Nil satis nisi optimum
CECILY: . . . Dr Chasuble is a most learned man. He has never written a single book, so you can imagine how much he knows.

Oscar Wilde

*The importance of being earnest*: 1899, Act III

The tendency to appear exact by disregarding the complexity of the factors is the old failing in our medical history.

J. Curnow

Cited in: Critchley M, Critchley EA.


FOREWORD TO THE FIRST EDITION

Neurology has always been a discipline in which careful physical examination is paramount. The rich vocabulary of neurology replete with eponyms attests to this historically. The decline in the importance of the examination has long been predicted with the advent of more detailed neuroimaging. However, neuroimaging has often provided a surfeit of information from which salient features have to be identified, dependent upon the neurological examination. A dictionary of neurological signs has a secure future.

A dictionary should be informative but unless it is unwieldy, it cannot be comprehensive, nor is that claimed here. Andrew Larner has decided sensibly to include key features of the history as well as the examination. There is no doubt that some features of the history can strike one with the force of a physical sign. There are entries for ‘palinopsia’ and ‘environmental tilt’ both of which can only be elicited from the history and yet which have considerable significance. There is also an entry for the ‘head turning sign’ observed during the history taking itself as well as the majority of entries relating to details of the physical examination.

This book is directed to students and will be valuable to medical students, trainee neurologists, and professions allied to medicine. Neurologists often speak in shorthand and so entries such as ‘absence’ and ‘freezing’ are sensible and helpful. For the more mature student, there are the less usual as well as common eponyms to entice one to read further than the entry which took you first to the dictionary.

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PREFACE TO THIRD EDITION

To paraphrase John Hughlings Jackson (1835–1911), clinical phenomena are experiments on the nervous system made by disease. Neurological signs might therefore be (loosely) characterized as the ‘dependent variables’ of the experiments wrought by neurological disease. Observing or eliciting these signs may therefore give insight into neurological disease processes. (Details of neurological disorders mentioned in passing in this book may be found elsewhere, in a companion volume.1) However, as mentioned in the preface to the first edition of this book,2 the accuracy and precision of most neurological signs remain to be defined at the level of traditional probabilistic parameters (specificity, sensitivity, positive and negative predictive values, likelihood ratios, etc.), perhaps because this undertaking is a less alluring prospect for research than, say, neuroimaging and neurogenetics. Thankfully, the clinical examination still has some supporters (not merely apologists), and neurological signs feature prominently amongst the core competencies.3 Clinicians, as opposed to academics, may be ideally placed to undertake the aforementioned studies in the context of their day-to-day clinical practice.

A.J. Larner

REFERENCES

ACKNOWLEDGEMENTS

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**Abadie’s Sign**
Abadie’s sign is the absence or diminution of pain sensation when exerting deep pressure on the Achilles tendon by squeezing. This is a frequent finding in the tabes dorsalis variant of neurosyphilis, i.e. with dorsal column disease.

**Cross Reference**
Argyll Robertson pupil

**Abdominal Paradox**
- see PARADOXICAL BREATHING

**Abdominal Reflexes**
Both superficial and deep abdominal reflexes are described, of which the superficial (cutaneous) reflexes are the more commonly tested in clinical practice. A wooden stick or pin is used to scratch the abdominal wall, from the flank to the midline, parallel to the line of the dermatomal strips, in upper (supraumbilical), middle (umbilical), and lower (infraumbilical) areas. The manoeuvre is best performed at the end of expiration when the abdominal muscles are relaxed, since the reflexes may be lost with muscle tensing; to avoid this, patients should lie supine with their arms by their sides.

  Superficial abdominal reflexes are lost in a number of circumstances:
  - Normal ageing;
  - Obesity;
  - Following abdominal surgery;
  - Following multiple pregnancies;
  - In acute abdominal disorders (Rosenbach’s sign).

  However, absence of all superficial abdominal reflexes may be of localizing value for corticospinal pathway damage (upper motor neurone lesions) above T6. Lesions at or below T10 lead to selective loss of the lower reflexes with the upper and middle reflexes intact, in which case Beevor’s sign may also be present. All abdominal reflexes are preserved with lesions below T12.

  Abdominal reflexes are said to be lost early in multiple sclerosis, but late in motor neurone disease, an observation of possible clinical use, particularly when differentiating the progressive lateral sclerosis variant of motor neurone disease from multiple sclerosis. However, no prospective study of abdominal reflexes in multiple sclerosis has been reported.

**Reference**

**Cross References**
Beevor’s sign; Upper motor neurone (UMN) syndrome
Abducens (VI) Nerve Palsy

Abducens (VI) nerve palsy causes a selective weakness of the lateral rectus muscle resulting in impaired abduction of the eye, manifest clinically as diplopia on lateral gaze, or on shifting gaze from a near to a distant object. Abducens (VI) nerve palsy may be due to:

- Microinfarction in the nerve, due to hypertension, diabetes mellitus;
- Raised intracranial pressure: a ‘false-localizing sign’, possibly caused by stretching of the nerve in its long intracranial course over the ridge of the petrous temporal bone;
- Nuclear pontine lesions: congenital, e.g. Duane retraction syndrome, Möbius syndrome;
- Unusual in multiple sclerosis.

Isolated weakness of the lateral rectus muscle may also occur in myasthenia gravis. In order not to overlook this fact, and miss a potentially treatable condition, it is probably better to label isolated abduction failure as ‘lateral rectus palsy’, rather than abducens nerve palsy, until the aetiological diagnosis is established.

Excessive or sustained convergence associated with a midbrain lesion (diencephalic–mesencephalic junction) may also result in slow or restricted abduction (pseudoabducens palsy, ‘midbrain pseudo-sixth’).

Reference


Cross References

Diplopia; ‘False-localizing signs’

Abductor Sign

The abductor sign is tested by asking the patient to abduct each leg whilst the examiner opposes movement with hands placed on the lateral surfaces of the patient’s legs: the leg contralateral to the abducted leg shows opposite actions dependent upon whether paresis is organic or non-organic. Abduction of a paretic leg is associated with the sound leg remaining fixed in organic paresis, but in non-organic paresis there is hyperadduction. Hence the abductor sign is suggested to be useful to detect non-organic paresis.

Reference


Cross Reference

Functional weakness and sensory disturbance

Absence

An absence, or absence attack, is a brief interruption of awareness of epileptic origin. This may be a barely noticeable suspension of speech or attentiveness, without postictal confusion or awareness that an attack has occurred, as in idiopathic generalized epilepsy of absence type (absence epilepsy; petit mal), a
Abulia

Abulia (aboulia) is a ‘syndrome of hypofunction’, characterized by a lack of initiative, spontaneity and drive (asplontaneity), apathy, slowness of thought (bradyphrenia), and blunting of emotional responses and response to external stimuli. It may be confused with the psychomotor retardation of depression and is sometimes labelled as ‘pseudodepression’. More plausibly, abulia has been thought of as a minor or partial form of akinetic mutism. A distinction may be drawn between abulia major (= akinetic mutism) and abulia minor, a lesser degree of abulia associated particularly with bilateral caudate stroke and thalamic infarcts in the territory of the polar artery and infratentorial stroke. There may also be some clinical overlap with catatonia.

Abulia may result from frontal lobe damage, most particularly that involving the frontal convexity, and has also been reported with focal lesions of the caudate nucleus, thalamus, and midbrain. As with akinetic mutism, it is likely that lesions anywhere in the ‘centromedial core’ of the brain, from frontal lobes to brainstem, may produce this picture.

Pathologically, abulia may be observed in:

- Infarcts in anterior cerebral artery territory and ruptured anterior communicating artery aneurysms, causing basal forebrain damage;
- Closed head injury;
- Parkinson’s disease; sometimes as a forerunner of a frontal lobe dementia;
- Other causes of frontal lobe disease: tumour, abscess;
- Metabolic, electrolyte disorders: hypoxia, hypoglycaemia, hepatic encephalopathy.

Treatment is of the underlying cause where possible. There is anecdotal evidence that the dopamine agonist bromocriptine may help.

References
Acalculia

Acalculia, or dyscalculia, is difficulty or inability in performing simple mental arithmetic. This depends on two processes, number processing and calculation; a deficit confined to the latter process is termed anarithmetia. Acalculia may be classified as:

- **Primary:**
  A specific deficit in arithmetical tasks, more severe than any other coexisting cognitive dysfunction.

- **Secondary:**
  In the context of other cognitive impairments, for example of language (aphasia, alexia, or agraphia for numbers), attention, memory, or space perception (e.g. neglect). Acalculia may occur in association with alexia, agraphia, finger agnosia, right–left disorientation, and difficulty spelling words as part of the Gerstmann syndrome with lesions of the dominant parietal lobe.

Secondary acalculia is the more common variety. Isolated acalculia may be seen with lesions of:

- dominant (left) parietal/temporal/occipital cortex, especially involving the angular gyrus (Brodmann areas 39 and 40);
- medial frontal lobe (impaired problem solving ability?);
- subcortical structures (caudate nucleus, putamen, internal capsule).

Impairments may be remarkably focal, for example one operation (e.g. subtraction) may be preserved whilst all others are impaired.

In patients with mild-to-moderate Alzheimer’s disease with dyscalculia but no attentional or language impairments, cerebral glucose metabolism was found to be impaired in the left inferior parietal lobule and inferior temporal gyrus. Preservation of calculation skills in the face of total language dissolution (production and comprehension) has been reported with focal left temporal lobe atrophy probably due to Pick’s disease.

References


Achromatopsia


Cross References
Agraphia; Alexia; Aphasia; Gerstmann syndrome; Neglect

Accommodation Reflex
- see PUPILLARY REFLEXES

Achilles Reflex

Plantar flexion at the ankle following phasic stretch of the Achilles tendon constitutes the Achilles reflex or ankle jerk, mediated through sacral segments S1 and S2 and the sciatic and posterior tibial nerves. This reflex may be elicited in several ways: by a blow with a tendon hammer directly upon the Achilles tendon (patient supine, prone with knee flexed, or kneeling) or with a plantar strike. The latter, though convenient and quick, is probably the least sensitive method, since absence of an observed muscle contraction does not mean that the reflex is absent; the latter methods are more sensitive.

The Achilles reflex is typically lost in polyneuropathies and S1 radiculopathy. Loss of the Achilles reflex is increasingly prevalent with normal healthy ageing, beyond the age of 60 years, although more than 65% of patients retain the ankle jerks.

References


Cross References
Age-related signs; Neuropathy; Reflexes

Achromatopsia

Achromatopsia, or dyschromatopsia, is an inability or impaired ability to perceive colours. This may be ophthalmological or neurological in origin, congenital or acquired; only in the latter case does the patient complain of impaired colour vision.
Achromatopsia is most conveniently tested for clinically using pseudoisochromatic figures (e.g. Ishihara plates), although these were specifically designed for detecting congenital colour blindness and test the red-green channel more than blue-yellow. Sorting colours according to hue, for example with the Farnsworth–Munsell 100 Hue test, is more quantitative, but more time-consuming. Difficulty performing these tests does not always reflect achromatopsia (see Pseudoachromatopsia).

Probably the most common cause of achromatopsia is inherited ‘colour blindness’, of which several types are recognized: in monochromats only one of the three cone photoreceptor classes is affected, in dichromats two; anomalous sensitivity to specific wavelengths of light may also occur (anomalous trichromat). These inherited dyschromatopsias are binocular, symmetrical, and do not change with time. Acquired achromatopsia may result from damage to the optic nerve or the cerebral cortex. Unlike inherited conditions, these deficits are noticeable (patients describe the world as looking ‘grey’ or ‘washed out’) and may be confined to only part of the visual field (e.g. hemiachromatopsia). Optic neuritis typically impairs colour vision (red-green > blue-yellow) and this defect may persist whilst other features of the acute inflammation (impaired visual acuity, central scotoma) remit. Cerebral achromatopsia results from cortical damage (most usually infarction) to the inferior occipitotemporal area. Area V4 of the visual cortex, which is devoted to colour processing, is in the occipitotemporal (fusiform) and lingual gyri. Unilateral lesions may produce a homonymous hemiachromatopsia. Lesions in this region may also produce prosopagnosia, alexia, and visual field defects, either a peripheral scotoma, which is always in the upper visual field, or a superior quadrantanopia, reflecting damage to the inferior limb of the calcarine sulcus in addition to the adjacent fusiform gyrus. Transient achromatopsia in the context of vertebrobasilar ischaemia has been reported.

The differential diagnosis of achromatopsia encompasses colour agnosia, a loss of colour knowledge despite intact perception; and colour anomia, an inability to name colours despite intact perception.

References

Cross References
Agnosia; Alexia; Anomia; Prosopagnosia; Pseudoachromatopsia; Quadrantanopia; Scotoma; Xanthopsia

Acousticopalpebral Reflex
- see BLINK REFLEX

Action Dystonia
- see DYSTONIA

Action Myoclonus
- see MYOCLONUS
Adiadochokinesia
- see DYSDIADOCHOKINESIA

Adie’s Syndrome, Adie’s Tonic Pupil
- see HOLMES–ADIE PUPIL, HOLMES–ADIE SYNDROME

Adson’s Test
Adson’s test may be helpful in the diagnosis of vascular thoracic outlet syndrome, along with Roos test. The arm is extended at the elbow, abducted, and then rotated posteriorly; following deep inspiration, the patient’s head is turned from one side to the other. Loss of the radial pulse may occur in normals but a bruit over the brachial artery is thought to suggest the presence of entrapment. A Doppler Adson’s test over the subclavian artery may predict successful outcome from thoracic outlet decompression surgery.

Reference

Cross Reference
Roos test

Adventitious Movements
- see STEREOTYPY

Affective Agnosia
- see AGNOSIA; APROSODIA, APROSODY

Afferent Pupillary Defect (APD)
- see RELATIVE AFFERENT PUPILLARY DEFECT (RAPD)

Age-Related Signs
A number of neurological signs are reported to be more prevalent with increasing age and related to ageing per se rather than any underlying age-related disease, hence not necessarily of pathological significance when assessing the neurological status of older individuals, although there are methodological difficulties in reaching such conclusions.

A brief topographical overview of age-related signs includes

- **Cognitive function:**
  Loss of processing speed, cognitive flexibility, efficiency of working memory (sustained attention);
  Preservation of vocabulary, remotely learned information including semantic networks, and well-encoded new information.

- **Cranial nerves:**
  *I*: olfactory sense diminished;
  *II, III, IV, VI*: presbyopia; reduced visual acuity, depth perception, contrast sensitivity, motion perception; ‘senile miosis’; restricted upward conjugate gaze;
Ageusia

VIII: presbycusis; impaired vestibulospinal reflexes.

- **Motor system:**
  
  *Appearance:* loss of muscle bulk; ‘senile’ tremor;
  
  *Tone:* rigidity; gegenhalten/paratonia;
  
  *Power:* decline in muscle strength;
  
  *Coordination:* impaired speed of movement (bradykinesia).

- **Reflexes:**
  
  *Phasic muscle stretch reflexes:* depressed or absent, especially ankle (Achilles tendon) jerk; jaw jerk;
  
  *Cutaneous (superficial) reflexes:* abdominal reflexes may be depressed with ageing;
  
  *Primitive/developmental reflexes:* glabellar, snout, palomemtal, grasp reflexes may be more common with ageing.

  Impairments of gait; parkinsonism.

- **Sensory system:**
  
  Decreased sensitivity to vibratory perception; +/- pain, temperature, proprioception

  Neuroanatomical correlates of some of these signs have been defined. There does seem to be an age-related loss of distal sensory axons and of spinal cord ventral horn motor neurones accounting for sensory loss, loss of muscle bulk and strength, and reflex diminution.

**References**


**Cross References**

Frontal release signs; Parkinsonism; Reflexes

**Ageusia**

Ageusia or hypogeusia is a loss or impairment of the sense of taste (gustation). This may be tested by application to each half of the protruded tongue the four fundamental tastes (sweet, sour, bitter, and salt).

Isolated ageusia is most commonly encountered as a transient feature associated with coryzal illnesses of the upper respiratory tract, as with anosmia. Indeed, many complaints of loss of taste are in fact due to anosmia, since olfactory sense is responsible for the discrimination of many flavours.

Neurological disorders may also account for ageusia. Afferent taste fibres run in the facial (VII) and glossopharyngeal (IX) cranial nerves, from taste buds
in the anterior two-thirds and posterior one-third of the tongue, respectively. Central processes run in the solitary tract in the brainstem and terminate in its nucleus (nucleus tractus solitarius), the rostral part of which is sometimes called the gustatory nucleus. Fibres then run to the ventral posterior nucleus of the thalamus, hence to the cortical area for taste adjacent to the general sensory area for the tongue (insular region).

Lesions of the facial nerve proximal to the departure of the chorda tympani branch in the mastoid (vertical) segment of the nerve (i.e. proximal to the emergence of the facial nerve from the stylomastoid foramen) can lead to ipsilateral impairment of taste sensation over the anterior two-thirds of the tongue, along with ipsilateral lower motor neurone facial weakness (e.g. in Bell’s palsy), with or without hyperacusis. Lesions of the glossopharyngeal nerve causing impaired taste over the posterior one-third of the tongue usually occur in association with ipsilateral lesions of the other lower cranial nerves (X, XI, XII; jugular foramen syndrome) and hence may be associated with dysphonia, dysphagia, depressed gag reflex, vocal cord paresis, anaesthesia of the soft palate, uvula, pharynx and larynx, and weakness of trapezius and sternocleidomastoid.

Ageusia as an isolated symptom of neurological disease is extremely rare, but has been described with focal central nervous system lesions (infarct, tumour, demyelination) affecting the nucleus of the tractus solitarius (gustatory nucleus) and/or thalamus and with bilateral insular lesions. Anosmia and dysgeusia have also been reported following acute zinc loss.

References

Cross References
Anosmia; Bell’s palsy; Cacogeusia; Dysgeusia; Facial paresis; Hyperacusis; Jugular foramen syndrome

Agnosia

Agnosia is a deficit of higher sensory (most often visual) processing causing impaired recognition. The term, coined by Freud in 1891, means literally ‘absence of knowledge’, but its precise clinical definition continues to be a subject of debate. Lissauer (1890) originally conceived of two kinds of agnosia:

- **Apperceptive:**
  In which there is a defect of complex (higher order) perceptual processes.

- **Associative:**
  In which perception is thought to be intact but there is a defect in giving meaning to the percept by linking its content with previously encoded percepts (the semantic system); this has been described as ‘a normal percept that has somehow been stripped of its meaning’ or ‘perception without knowledge’.

- 9 -
These deficits should not be explicable by a concurrent intellectual impairment, disorder of attention, or by an inability to name or describe verbally the stimulus (anomia). As a corollary of this last point, some argue that there should be no language disorder (aphasia) to permit the diagnosis of agnosia.

Intact perception is sometimes used as a sine qua non for the diagnosis of agnosia, in which case it may be questioned whether apperceptive agnosia is truly agnosia. However, others retain this category, not least because the supposition that perception is normal in associative visual agnosia is probably not true. Moreover, the possibility that some agnosias are in fact higher-order perceptual deficits remains: examples include some types of visual and tactile recognition of form or shape (e.g. agraphognosia; astereognosis; dysmorphopsia); some authorities label these phenomena as ‘pseudoagnosias’. The difficulty with definition perhaps reflects the continuing problem of defining perception at the physiological level. Other terms which might replace agnosia have been suggested, such as non-commitittal terms like ‘disorder of perception’ or ‘perceptual defect’, or as suggested by Hughlings Jackson ‘imperception’.

Theoretically, agnosias can occur in any sensory modality, but some authorities believe that the only unequivocal examples are in the visual and auditory domains (e.g. prosopagnosia and pure word deafness, respectively). Nonetheless, many other ‘agnosias’ have been described, although their clinical definition may lie outwith some operational criteria for agnosia. With the passage of time, agnostic defects merge into anterograde amnesia (failure to learn new information).

Anatomically, agnosias generally reflect dysfunction at the level of the association cortex, although they can on occasion result from thalamic pathology. Some may be of localizing value. The neuropsychological mechanisms underpinning these phenomena are often ill understood.

References

Cross References
Agraphognosia; Alexia; Amnesia; Anosognosia; Aprosodia, Aprosody; Asomatognosia; Astereognosia; Auditory agnosia; Autotopagnosia; D ysmorphopsia; Finger agnosia; Phonagnosia; Prosopagnosia; Pure word deafness; Simultanagnosia; Tactile agnosia; Visual agnosia; Visual form agnosia

Agrammatism
Agrammatism is a reduction in, or loss of, the production or comprehension of the syntactic elements of language, for example articles, prepositions, conjunctions, verb endings (i.e. the non-substantive components of language), whereas nouns and verbs are relatively spared. Despite this impoverishment of language,
Agraphia or ‘telegraphic speech’, meaning is often still conveyed because of the high information content of verbs and nouns. Agrammatism is encountered in Broca’s type of non-fluent aphasia, associated with lesions of the posterior inferior part of the frontal lobe of the dominant hemisphere (Broca’s area). Agrammatic speech may also be dysprosodic.

**Cross References**

Aphasia; Aprosodia, Aprosody

### Agraphaesthesia

Agraphaesthesia, dysgraphaesthesia, or graphanaesthesia is a loss or impairment of the ability to recognize letters or numbers traced on the skin, i.e. of graphaesthesia. Whether this is a perceptual deficit or a tactile agnosia (‘agraphognosia’) remains a subject of debate. It occurs with damage to the somatosensory parietal cortex. It may occur in corticobasal degeneration syndrome.

**Cross References**

Agnosia; Tactile agnosia

### Agraphia

Agraphia or dysgraphia is a loss or disturbance of the ability to write or spell. Since writing depends not only on language function but also on motor, visuospatial, and kinaesthetic function, many factors may lead to dysfunction. Agraphias may be classified as follows:

- **Central, aphasic, or linguistic dysgraphias:**
  
  These are usually associated with aphasia and alexia, and the deficits mirror those seen in the Broca/anterior/motor and Wernicke/posterior/sensory types of aphasia. Oral spelling is impaired. From the linguistic viewpoint, two types of paragraphia may be distinguished as follows:

  - **Surface/lexical/semantic dysgraphia:** misspelling of irregular words, producing phonologically plausible errors (e.g. simtums for symptoms); this is seen with left temporoparietal lesions, e.g. Alzheimer’s disease, Pick’s disease;
  
  - **Deep/phonological dysgraphia:** inability to spell unfamiliar words and non-words; semantic errors; seen with extensive left hemisphere damage.

- **Mechanical agraphia:**

  Impaired motor control, due to paresis (as in dominant parietal damage), dyspraxia (may be accompanied by ideomotor limb apraxia), dyskinesia (hypokinetic or hyperkinetic), or dystonia; oral spelling may be spared.

- **Neglect (spatial) dysgraphia:**

  Associated with other neglect phenomena consequent upon a non-dominant hemisphere lesion; there may be missing out or misspelling of the left side of words (paragraphia); oral spelling may be spared.

- **Pure agraphia:**

  A rare syndrome in which oral language, reading, and praxis are normal.
A syndrome of agraphia, alexia, acalculia, finger agnosia, right–left disorientation, and difficulty spelling words (Gerstmann syndrome) may be seen with dominant parietal lobe pathologies.

Writing disturbance due to abnormal mechanics of writing is the most sensitive language abnormality in delirium, possibly because of its dependence on multiple functions.

References

Cross References
Alexia; Allographia; Aphasia; Apraxia; Broca’s aphasias; Fast micrographia; Gerstmann syndrome; Hypergraphia; Macrographia; Micrographia; Neglect; Wernicke’s aphasia

Agraphognosia
- see AGRAPHAESTHESIA

Agrypnia
Agrypnia, or agrypnia excitata, is severe, total insomnia of long duration. Recognized causes include trauma to the brainstem and/or thalamus, prion disease (fatal familial and sporadic fatal insomnia), Morvan’s syndrome, von Economo’s disease, trypanosomiasis, and a relapsing-remitting disorder of possible autoimmune pathogenesis responding to plasma exchange.

References

Akathisia
Akathisia is a feeling of inner restlessness, often associated with restless movements of a continuous and often purposeless nature, such as rocking to and fro, repeatedly crossing and uncrossing the legs, standing up and sitting down, and pacing up and down (forced walking, tasikinesia). Moaning, humming, and groaning may also be features. Voluntary suppression of the movements may exacerbate inner tension or anxiety.

Recognized associations of akathisia include Parkinson’s disease and neuroleptic medication use (acute or tardive side effect), suggesting that dopamine depletion may contribute to the pathophysiology. Dopamine-depleting agents (e.g. tetrabenazine, reserpine) may also cause akathisia. The Barnes Akathisia Rating Scale is the standard assessment scale.

Treatment of akathisia by reduction or cessation of neuroleptic therapy may help, but may exacerbate coexistent psychosis. Centrally acting β-blockers such as propranolol may also be helpful, as may anticholinergic agents, amantadine, clonazepam, and clonidine.
Akinesia

Akinesia is a lack of, or an inability to initiate, voluntary movements. More usually in clinical practice there is a difficulty (reduction, delay), rather than complete inability, in the initiation of voluntary movement, perhaps better termed bradykinesia, or reduced amplitude of movement or hypokinesia. These difficulties cannot be attributed to motor unit or pyramidal system dysfunction. Reflexive motor activity may be preserved (kinesis paradoxica). There may be concurrent slowness of movement, also termed bradykinesia.

Akinesia may coexist with any of the other clinical features of extrapyramidal system disease, particularly rigidity, but the presence of akinesia is regarded as an absolute requirement for the diagnosis of parkinsonism. Hemiakinesia may be a feature of motor neglect of one side of the body (possibly a motor equivalent of sensory extinction). Bilateral akinesia with mutism (akinetic mutism) may occur if pathology is bilateral. Pure akinesia, without rigidity or tremor, may occur: if levodopa-responsive, this is usually due to Parkinson’s disease; if levodopa-unresponsive, it may be the harbinger of progressive supranuclear palsy. A few patients with PSP have ‘pure akinesia’ without other features until late in the disease course.

Neuroanatomically, akinesia is a feature of disorders affecting

- frontal–subcortical structures, e.g. the medial convexity subtype of frontal lobe syndrome;
- basal ganglia;
- ventral thalamus;
- limbic system (anterior cingulate gyrus).

Neurophysiologically, akinesia is associated with loss of dopamine projections from the substantia nigra to the putamen.

Pathological processes underpinning akinesia include

- Neurodegeneration, e.g. Parkinson’s disease, progressive supranuclear palsy (Steele–Richardson–Olszewski syndrome), and multiple system atrophy (striatonigral degeneration); akinesia may occur in frontotemporal lobar degeneration syndromes, Alzheimer’s disease, and some prion diseases;
- Hydrocephalus;
- Neoplasia, e.g. butterfly glioma of the frontal lobes;
- Cerebrovascular disease.

Akinesia resulting from nigrostriatal dopamine depletion (i.e. idiopathic Parkinson’s disease) may respond to treatment with levodopa or dopamine agonists. However, many parkinsonian/akinetic-rigid syndromes show no or only partial response to these agents.

References

Cross References
Parkinsonism; Tasikinesia; Tic
Akinetic Mutism

Akinetic mutism is a ‘syndrome of negatives’, characterized by a lack of voluntary movement (akinesia), absence of speech (mutism), and lack of response to question and command, but with normal alertness and sleep–wake cycles (cf. coma). Blinking (spontaneous and to threat) is preserved. Frontal release signs, such as grasping and sucking, may be present, as may double incontinence, but there is a relative paucity of upper motor neurone signs affecting either side of the body, suggesting relatively preserved descending pathways. Akinetic mutism represents an extreme form of abulia, hence sometimes referred to as abulia major.

Pathologically, akinetic mutism is associated with bilateral lesions of the ‘centromedial core’ of the brain interrupting reticular-cortical or limbic-cortical pathways but which spare corticospinal pathways; this may occur at any point from frontal lobes to brainstem. Two forms of akinetic mutism are sometimes distinguished:

- **Frontodiencephalic**: associated with bilateral occlusion of the anterior cerebral arteries or with haemorrhage and vasospasm from anterior communicating artery aneurysms; damage to the cingulate gyri appears crucial but not sufficient for this syndrome.
- **Akinetic mutism with disturbances of vertical eye movements and hypersomnia**: associated with paramedian thalamic and thalamomesencephalic strokes.

Other structures (e.g. globus pallidus) have sometimes been implicated. Pathology may be vascular, neoplastic, or structural (subacute communicating hydrocephalus), and evident on structural brain imaging. Akinetic mutism may be the final state common to the end-stages of a number of neurodegenerative pathologies. EEG may show slowing with a lack of desynchronization following external stimuli.

Occasionally, treatment of the cause may improve akinetic mutism (e.g. relieving hydrocephalus). Agents such as dopamine agonists (e.g. bromocriptine), ephedrine and methylphenidate have also been tried.

References


Cross References
Abulia; Akinesia; Athymhormia; Blink reflex; Catatonia; Coma; Frontal lobe syndromes; Frontal release signs; Grasp reflex; Locked-in syndrome; Mutism

Akinetic-Rigid Syndrome
- see PARKINSONISM

Akinetopsia
Akinetopsia is a specific inability to see objects in motion, the perception of other visual attributes, such as colour, form, and depth, remaining intact. This statokinetic dissociation may be known as Riddoch’s phenomenon; the syndrome may also be called cerebral visual motion blindness. Such cases, although exceptionally rare, suggest a distinct neuroanatomical substrate for movement vision, as do cases in which motion vision is selectively spared in a scotomatous area (Riddoch’s syndrome).

Akinetopsia reflects a lesion selective to area V5 of the visual cortex. Clinically, it may be associated with acalculia and aphasia.

References

Cross References
Acalculia; Aphasia; Riddoch’s phenomenon

Alalia
Alalia is now an obsolete term, once used to describe a disorder of the material transformation of ideas into sounds. Lordat used it to describe the aphasia following a stroke.

Reference

Cross References
Aphasia; Aphemia

Alexia
Alexia is an acquired disorder of reading. The word dyslexia, though in some ways equivalent, is often used to denote a range of disorders in people who fail to develop normal reading skills in childhood. Alexia may be described as an acquired dyslexia. Alexia may be categorized as:

- Peripheral:
  A defect of perception or decoding the visual stimulus (written script); other language functions are often intact.
Peripheral alexias include

- **Alexia without agraphia:**
  Also known as pure alexia or pure word blindness. This is the archetypal peripheral alexia. Patients lose the ability to recognize written words quickly and easily; they seem unable to process all the elements of a written word in parallel. They can still access meaning but adopt a laborious letter-by-letter strategy for reading, with a marked word-length effect (i.e. greater difficulty reading longer words). Patients with pure alexia may be able to identify and name individual letters, but some cannot manage even this (‘global alexia’). Strikingly the patient can write at normal speed (i.e. no agraphia) but is then unable to read what they have just written. Alexia without agraphia often coexists with a right homonymous hemianopia, and colour anoma or impaired colour perception (achromatopsia); this latter may be restricted to one hemifield, classically right-sided (hemichromatopsia). Pure alexia has been characterized by some authors as a limited form of associative visual agnosia or ventral simultanagnosia.

- **Hemianopic alexia:**
  This occurs when a right homonymous hemianopia encroaches into central vision. Patients tend to be slower with text than single words as they cannot plan rightward reading saccades.

- **Neglect alexia:**
  Or hemiparalexia, results from failure to read either the beginning or end of a word (more commonly the former) in the absence of a hemianopia, due to hemispatial neglect.

The various forms of peripheral alexia may coexist; following a stroke, patients may present with global alexia which evolves to a pure alexia over the following weeks. Pure alexia is caused by damage to the left occipitotemporal junction, its afferents from early mesial visual areas, or its efferents to the medial temporal lobe. Global alexia usually occurs when there is additional damage to the splenium or white matter above the occipital horn of the lateral ventricle. Hemianopic alexia is usually associated with infarction in the territory of the posterior cerebral artery damaging geniculo striate fibres or area V1 itself, but can be caused by any lesion outside the occipital lobe that causes a macular splitting homonymous field defect. Neglect alexia is usually caused by occipitoparietal lesions, right-sided lesions causing left neglect alexia.

Central (linguistic) alexias include

- **Alexia with aphasia:**
  Patients with aphasia often have coexistent difficulties with reading (reading aloud and/or comprehending written text) and writing (alexia with agraphia, such patients may have a complete or partial Gerstmann
syndrome, the so-called third alexia of Benson). The reading problem parallels the language problem; thus in Broca’s aphasia reading is laboured with particular problems in reading function words (of, at) and verb inflections (-ing, -ed); in Wernicke’s aphasia numerous paraphasic errors are made.

From the linguistic viewpoint, different types of paralexia (substitution in reading) may be distinguished:

- **Surface dyslexia:**
  
  *Reading by sound*: there are regularization errors with exception words (e.g. pint pronounced to rhyme with mint), but non-words can be read; this may be seen with left medial +/- lateral temporal lobe pathology, e.g. infarction, semantic dementia, and late Alzheimer’s disease.

- **Phonological dyslexia:**
  
  *Reading by sight*: difficulties with suffixes, unable to read non-words; left temporoparietal lobe pathology.

- **Deep dyslexia:**
  
  The inability to translate orthography to phonology, manifesting as an inability to read plausible non-words (as in phonological dyslexia), plus semantic errors related to word meaning rather than sound (e.g. sister read as uncle); visual errors are also common (e.g. sacred read as scared). Deep dyslexia is seen with extensive left hemisphere temporoparietal damage.

The term transcortical alexia has been used to describe patients with Alzheimer’s disease with severe comprehension deficits who nonetheless are able to read aloud virtually without error all regular and exception words.

**References**


**Cross References**

Acalculia; Achromatopsia; Agnosia; Agraphia; Aphasia; Broca’s aphasia; Gerstmann syndrome; Hemianopia; Macula sparing, Macula splitting; Neglect; Prosopagnosia; Saccades; Simultanagnosia; Visual agnosia; Visual field defects; Wernicke’s aphasia

**Alexithymia**

Alexithymia is a reduced ability to identify and express ones feelings. This may contribute to various physical and behavioural disorders. It may be measured...
using the Toronto Alexithymia Score. There is evidence from functional imaging studies that alexithymics process facial expressions differently from normals, leading to the suggestion that this contributes to disordered affect regulation. Alexithymia is a common finding in split-brain patients, perhaps resulting from disconnection of the hemispheres.

**References**

‘Alice in Wonderland’ Syndrome

The name ‘Alice in Wonderland’ syndrome was coined by Todd in 1955 to describe the phenomena of microsomatognosia or macrosomatognosia, altered perceptions of body image, although these had first been described by Lippman in the context of migraine some years earlier. It has subsequently been suggested that Charles Lutwidge Dodgson’s own experience of migraine, recorded in his diaries, may have given rise to Lewis Carroll’s descriptions of Alice’s changes in body form, graphically illustrated in *Alice’s Adventures in Wonderland* (1865) by Sir John Tenniel. Some authors have subsequently interpreted these as somesthetic migrainous auras whereas others challenge this on chronological grounds, finding no evidence in Dodgson’s diaries for the onset of migraine until after he had written the Alice books. Moreover, migraine with somesthetic auras is rare, and Dodgson’s diaries have no report of migraine-associated body image hallucinations.

Other conditions may also give rise to the phenomena of microsomatognosia or macrosomatognosia, including epilepsy, encephalitis, cerebral mass lesions, schizophrenia, and drug intoxication.

**References**

**Cross References**
Aura; Metamorphopsia.

Alien Grasp Reflex

The term alien grasp reflex has been used to describe a grasp reflex occurring in full consciousness, which the patient could anticipate but perceived as alien (i.e. not modified by will), occurring in the absence of other abnormal movements. These phenomena were associated with an intrinsic tumour of the right (non-dominant) frontal lobe. It was suggested that the grasp reflex and alien hand syndromes are not separate entities but part of the spectrum of frontal lobe dysfunction, the term ‘alien grasp reflex’ attempting to emphasize the overlap.
Alien Hand, Alien Limb

An alien limb, most usually the arm but occasionally the leg, is one which manifests slow, involuntary, wandering (levitating), quasi-purposive movements. An arm so affected may show apraxic difficulties in performing even the simplest tasks and may be described by the patient as uncooperative or ‘having a mind of its own’ (hence alternative names such as anarchic hand sign, le main étranger, and ‘Dr Strangelove syndrome’). These phenomena are often associated with a prominent grasp reflex, forced groping, intermanual conflict, and magnetic movements of the hand. Different types of alien hand have been described, reflecting the differing anatomical locations of underlying lesions:

- **Anterior or motor types:**
  - *Callosal type:* characterized primarily by intermanual conflict.
  - *Frontal type:* shows features of environmental dependency, such as forced grasping and groping, and utilization behaviour.
- **Sensory or posterior variant:**
  - Resulting from a combination of cerebellar, optic, and sensory ataxia; rare.

A paroxysmal alien hand has been described, probably related to seizures of frontomedial origin.

Recognized pathological associations of alien limb include

- Corticobasal (ganglionic) degeneration;
- Corpus callosum tumours, haemorrhage;
- Medial frontal cortex infarction (territory of the anterior cerebral artery);
- Trauma and haemorrhage affecting both corpus callosum and medial frontal area;
- Alzheimer’s disease, familial Creutzfeldt–Jakob disease (very rare);
- Posterior cerebral artery occlusion (sensory variant);
- Following commissurotomy (corpus callosotomy alone insufficient).

Functional imaging studies in corticobasal degeneration, along with the evidence from focal vascular lesions, suggest that damage to and/or hypometabolism of the medial frontal cortex (Brodmann area 32) and the supplementary motor area (Brodmann area 6) is associated with alien limb phenomena. More generally, it seems that these areas are involved in the execution of learned motor programs, and damage thereto may lead to the release of learned motor programs from voluntary control.

References


**Cross References**

Alien grasp reflex; Apraxia; Ataxia; ‘Compulsive grasping hand’; Forced groping; Grasp reflex; Intermanual conflict; Levitation; Magnetic movements; Utilization behaviour

**Alienation Du Mot**

A loss of the feeling of familiarity with a word, part of the comprehension deficit seen in semantic dementia.

**Reference**


**Alloacousia**

Alloacousia describes a form of auditory neglect seen in patients with unilateral spatial neglect, characterized by spontaneous ignoring of people addressing the patient from the contralesional side, failing to respond to questions, or answering as if the speaker were on the ipsilesional side.

**Reference**


**Cross Reference**

Neglect

**Alloaesthesia**

Alloaesthesia (allesthesia, alloesthesia) is the condition in which a sensory stimulus given to one side of the body is perceived at the corresponding area on the other side of the body after a delay of about half a second. The trunk and proximal limbs are affected more often than the face or distal limbs. Visual alloaesthesia, the illusory transposition of an object seen in one visual field to the contralateral visual field, is also described, for example in ‘top of the basilar’ syndrome or with occipital lobe tumours. Tactile alloaesthesia may be seen in the acute stage of right putaminal haemorrhage (but seldom in right thalamic haemorrhage) and occasionally with anterolateral spinal cord lesions. The author has seen a patient report sensation below the stump of an amputated leg following stimulation of the contralateral remaining leg, a phenomenon which might be termed ‘phantom alloaesthesia’. ‘Mirror pain’, which has been reported after percutaneous cordotomy interrupting spinothalamic tracts to alleviate refractory pain syndromes (ML Sharma, personal communication), may share a similar neurobiological substrate. The mechanism of alloaesthesia is uncertain: some
consider it a disturbance within sensory pathways, others consider that it is a sensory response to neglect.

**References**

**Cross References**
Allochiria; Allokinesia, Allokinesis; Neglect

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**Allochiria**

Allochiria is the mislocation of sensory stimuli to the corresponding half of the body or space, a term coined by Obersteiner in 1882. There is overlap with alloaesthesia, originally used by Stewart (1894) to describe stimuli displaced to a different point on the same extremity.

Transposition of objects may occur in patients with neglect, e.g. from the neglected side (usually left) to the opposite side (usually right): for example, in a patient with left visuospatial neglect from a right frontoparietal haemorrhage, a figure was copied with objects from the left side transposed to the right.

**References**

**Cross References**
Alloaesthesia; Allokinesia, Allokinesis; Neglect; Right–left disorientation

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**Allodynia**

Allodynia is the elicitation of pain by light mechanical stimuli (such as touch or light pressure) which do not normally provoke pain (cf. hyperalgesia), i.e. this is a positive sensory phenomenon. Examples of allodynia include the trigger points of trigeminal neuralgia, the affected skin in areas of causalgia, and some peripheral neuropathies; it may also be provoked, paradoxically, by prolonged morphine use.

Various pathogenetic mechanisms are considered possible, including sensitization (lower threshold, hyperexcitability) of peripheral cutaneous nociceptive fibres (in which neurotrophins may play a role); ephaptic transmission (‘cross-talk’) between large and small (nociceptive) afferent fibres; and abnormal central processing.

The treatment of neuropathic pain is typically with agents such as carbamazepine, amitriptyline, gabapentin, and pregabalin. Interruption of sympathetic outflow, for example with regional guanethidine blocks, may sometimes help, but relapse may occur.

**Cross References**
Hyperalgesia; Hyperpathia
Allographia
This term has been used to describe a peripheral agraphia syndrome characterized by problems spelling both words and non-words, with case change errors such that upper and lower case letters are mixed when writing, with upper and lower case versions of the same letter sometimes superimposed on one another. Such errors increased in frequency with word length. These defects have been interpreted as a disturbance in selection of allographic forms in response to graphemic information outputted from the graphemic response buffer.

Reference

Cross Reference
Agraphia

Allokinesia, Allokinesis
Allokinesia has been used to denote a motor response in the wrong limb (e.g. movement of the left leg when attempting to move a paretic left arm) or transposition of the intended movement to the contralateral side; the movement may also be in the wrong direction. Others have used the term to denote a form of motor neglect, akin to alloaesthesia and allochiria in the sensory domain, relating to incorrect responses in the limb ipsilateral to a frontal lesion, also labelled disinhibition hyperkinesia.

References

Cross References
Alloaesthesia; Allochiria; Neglect

Alternate Cover Test
- see COVER TESTS

Alternating Fist Closure Test
In the alternating fist closure test, patients are asked to open and close the fists alternating (i.e. open left, close right, and vice versa) at a comfortable rate. Patients with limb-kinetic apraxia cannot keep pace and lose track.

Cross References
Apraxia; Frontal lobe syndromes

Alternating Sequences Test
- see APRAXIA; FRONTAL LOBE SYNDROMES

Altitudinal Field Defect
Altitudinal visual field defects are horizontal hemianopsias, in that they respect the horizontal meridian; they may be superior or inferior. Altitudinal field defects
Amblyopia are characteristic of (but not exclusive to) disease in the distribution of the central retinal artery. Central vision may be preserved (macula sparing) because the blood supply of the macula often comes from the cilioretinal arteries. Recognized causes of altitudinal visual field defects include

- **Monocular:**
  - Central retinal artery occlusion (CRAO);
  - Acute ischaemic optic neuropathy (AION);
  - Retinal detachment;
  - Choroiditis;
  - Glaucoma;
  - Chronic atrophic papilloedema.

- **Bilateral:**
  - Sequential CRAO, AION;
  - Bilateral occipital (inferior or superior calcarine cortices) lesions.

**Cross References**
Hemianopia; Macula sparing, Macula splitting; Quadrantanopia; Visual field defects

**Amaurosis**
Amaurosis is visual loss, with the implication that this is not due to refractive error or intrinsic ocular disease. The term is most often used in the context of amaurosis fugax, a transient monocular blindness, which is most often due to embolism from a stenotic ipsilateral internal carotid artery (ocular transient ischaemic attack). Giant cell arteritis, systemic lupus erythematosus, and the antiphospholipid antibody syndrome are also recognized causes. Gaze-evoked amaurosis has been associated with a variety of mass lesions and is thought to result from decreased blood flow to the retina from compression of the central retinal artery with eye movement.

**Amblyopia**
Amblyopia refers to poor visual acuity, most usually in the context of a ‘lazy eye’, in which the poor acuity results from the failure of the eye to establish normal cortical representation of visual input during the critical period of visual maturation (between the ages of 6 months and 3 years). This may result from:

- strabismus;
- uncorrected refractive error;
- stimulus deprivation.

Amblyopic eyes may demonstrate a relative afferent pupillary defect and sometimes latent nystagmus.

Amblyopia may not become apparent until adulthood, when the patient suddenly becomes aware of unilateral poor vision. The finding of a latent strabismus (heterophoria) may be a clue to the fact that such visual loss is long-standing.

The word amblyopia has also been used in other contexts: bilateral simultaneous development of central or centrocaecal scotomas in chronic alcoholics has often been referred to as tobacco–alcohol amblyopia, although nutritional optic neuropathy is perhaps a better term.
Amnesia

Amnesia is an impairment of episodic memory or memory for personally experienced events (autobiographical memory). This is a component of long-term (as opposed to working) memory which is distinct from memory for facts (semantic memory), in that episodic memory is unique to the individual whereas semantic memory encompasses knowledge held in common by members of a cultural or linguistic group. Episodic memory generally accords with the lay perception of memory, although many complaints of ‘poor memory’ represent faulty attentional mechanisms rather than true amnesia. A precise clinical definition for amnesia has not been demarcated, perhaps reflecting the heterogeneity of the syndrome.

Amnesia may be retrograde (for events already experienced) or anterograde (for newly experienced events). Retrograde amnesia may show a temporal gradient, with distant events being better recalled than more recent ones, relating to the duration of anterograde amnesia.

Amnesia may be acute and transient or chronic and persistent. In a pure amnesic syndrome, intelligence and attention are normal and skill acquisition (procedural memory) is preserved. Amnesia may occur as one feature of more widespread cognitive impairments, e.g. in Alzheimer’s disease.

Various psychometric tests of episodic memory are available. These include the Wechsler Memory Score-Revised (WMS-R), the Recognition Memory Test which has both verbal (words) and visual (faces) subdivisions, the Rey Auditory Verbal Learning Test (immediate and delayed free recall of a random word list), and the Rey–Osterreith Complex Figure (non-verbal memory). Retrograde memory may be assessed with a structured Autobiographical Memory Interview and with the Famous Faces Test. Poor spontaneous recall, for example, of a word list, despite an adequate learning curve, may be due to a defect in either storage or retrieval. This may be further probed with cues: if this improves recall, then a disorder of retrieval is responsible; if cueing leads to no improvement or false-positive responses to foils (as in the Hopkins Verbal Learning Test) are equal or greater than true positives, then a learning defect (true amnesia) is the cause.

The neuroanatomical substrate of episodic memory is a distributed system in the medial temporal lobe and diencephalon surrounding the third ventricle (the circuit of Papez) comprising the entorhinal area of the parahippocampal gyrus, perforant and alvear pathways, hippocampus, fimbria and fornix, mammillary bodies, mammillothalamic tract, anterior thalamic nuclei, internal capsule, cingulate gyrus, and cingulum. Basal forebrain structures (septal nucleus, diagonal band nucleus of Broca, nucleus basalis of Meynert) are also involved.
Classification of amnesic syndromes into subtypes has been proposed, since lesions in different areas produce different deficits reflecting functional subdivision within the system; thus left temporal lesions produce problems in the verbal domain, right-sided lesions affect non-verbal/visual memory. A distinction between medial temporal pathology (e.g. hippocampus), leading to difficulty encoding new memories (anterograde amnesia and temporally limited retrograde amnesia), and diencephalic pathology (e.g. Korsakoff’s syndrome), which causes difficulty retrieving previously acquired memories (extensive retrograde amnesia) with diminished insight and a tendency to confabulation, has been suggested, but overlap may occur. A frontal amnesia has also been suggested, although impaired attentional mechanisms may contribute. Functional imaging studies suggest that medial temporal lobe activation is required for encoding with additional prefrontal activation with ‘deep’ processing; medial temporal and prefrontal activations are also seen with retrieval.

Many causes of amnesia are recognized, including

- **Acute/transient:**
  - Closed head injury;
  - Drugs;
  - Transient global amnesia;
  - Transient epileptic amnesia;
  - Transient semantic amnesia (very rare).

- **Chronic/persistent:**
  - Alzheimer’s disease (may show isolated amnesia in early disease);
  - Sequela of herpes simplex encephalitis;
  - Limbic encephalitis (paraneoplastic or non-paraneoplastic);
  - Hypoxic brain injury;
  - Temporal lobectomy (bilateral; or unilateral with previous contralateral injury, usually birth asphyxia);
  - Bilateral posterior cerebral artery occlusion;
  - Korsakoff’s syndrome;
  - Bilateral thalamic infarction;
  - Third ventricle tumour, cyst;
  - Focal retrograde amnesia (rare).

Few of the chronic persistent causes of amnesia are amenable to specific treatment. Plasma exchange or intravenous immunoglobulin therapy may be helpful in non-paraneoplastic limbic encephalitis associated with autoantibodies directed against voltage-gated potassium channels.

Functional or psychogenic amnesia may involve failure to recall basic autobiographical details such as name and address. Reversal of the usual temporal gradient of memory loss may be observed (but this may also be the case in the syndrome of focal retrograde amnesia).

**References**


Amphigory


**Cross References**
Confabulation; Dementia; Dissociation

**Amphigory**
Fisher used this term to describe nonsense speech.

**References**

**Cross Reference**
Aphasia

**Amusia**
Amusia is a loss of the ability to appreciate music despite normal intelligence, memory, and language function. Subtypes have been described: receptive or sensory amusia is loss of the ability to appreciate music; and expressive or motor amusia is loss of ability to sing, whistle. Clearly a premorbid appreciation of music is a sine qua non for the diagnosis (particularly of the former), and most reported cases of amusia have occurred in trained musicians. Others have estimated that amusia affects up to 4% of the population (presumably expressive; = ‘tone deafness’). Tests for the evaluation of amusia have been described.

Amusia may occur in the context of more widespread cognitive dysfunction, such as aphasia and agnosia. It has been found in association with pure word deafness, presumably as part of a global auditory agnosia. Isolated amusia has been reported in the context of focal cerebral atrophy affecting the non-dominant temporal lobe. However, functional studies have failed to show strong hemispheric specificity for music perception, but suggest a cross-hemispheric distributed neural substrate. An impairment of pitch processing with preserved awareness of musical rhythm changes has been described in amusics.

**References**

**Cross References**
Agnosia; Auditory agnosia; Pure word deafness
Amyotrophy
Amyotrophy is a term used to describe thinning or wasting (atrophy) of musculature with attendant weakness. This may result from involvement of:

- **Lower motor neurones** (in which case fasciculations may also be present):
  - Amyotrophic lateral sclerosis/motor neurone disease;
  - Benign focal amyotrophy/monomelic amyotrophy;
  - Disinhibition-dementia-parkinsonism-amyotrophy complex (DDPAC);
  - ‘Asthmatic amyotrophy’ (Hopkins’ syndrome).

- **Nerve roots**:
  - Diabetic amyotrophy (polyradiculopathy, especially L2–L4).

- **Plexus**:
  - Neuralgic amyotrophy (Parsonage–Turner syndrome).

Hence, although the term implies neurogenic (as opposed to myogenic) muscle wasting, its use is non-specific with respect to neuroanatomical substrate.

Cross References
Atrophy; Fasciculation; Neuropathy; Plexopathy; Radiculopathy; Wasting

Anaesthesia
Anaesthesia (anesthesia) is a complete loss of sensation; hypoaesthesia (hypesthesia) is a diminution of sensation. Hence in Jacksonian terms, these are negative sensory phenomena. Anaesthesia may involve all sensory modalities (global anaesthesia, as in general surgical anaesthesia) or be selective (e.g. thermoanaesthesia, analgesia). Regional patterns of anaesthesia are described, e.g. ‘glove-and-stocking anaesthesia’ in peripheral neuropathies and ‘saddle anaesthesia’ involving S3-5 dermatomes resulting from a cauda equina syndrome.

Anaesthesia is most often encountered after resection or lysis of a peripheral nerve segment, whereas paraesthesia or dysaesthesia (positive sensory phenomena) reflects damage to a nerve which is still in contact with the cell body.

Anaesthesia dolorosa, or painful anaesthesia, is a persistent unpleasant pain (i.e. a positive sensory phenomenon) which may be experienced in the distribution of a resected nerve, e.g. following neurolytic treatment for trigeminal neuralgia, usually with delayed onset. This deafferentation pain may respond to various medications, including tricyclic antidepressants, carbamazepine, gabapentin, pregabalin, and selective serotonin-reuptake inhibitors.

Cross References
Analgesia; Dysesthesia; Neuropathy; Paraesthesia

Analgesia
Analgesia or hypoalgesia refers to a complete loss or diminution, respectively, of pain sensation, or the absence of a pain response to a normally painful stimulus. These negative sensory phenomena may occur as one component of total sensory loss (anaesthesia) or in isolation. Consequences of analgesia include
the development of neuropathic ulcers, burns, Charcot joints, even painless mutilation, or amputation. Analgesia may occur in:

- peripheral nerve lesions, e.g. hereditary sensory and autonomic neuropathies (HSAN), leprosy;
- central spinal cord lesions which pick off the decussating fibres of the spinothalamic pathway in the ventral funiculus (with corresponding thermoanaesthesia), e.g. syringomyelia;
- cortical lesions, e.g. medial frontal lobe syndrome (akinetic type).

Congenital syndromes of insensitivity to pain were once regarded as a central pain asymbolia (e.g. Osuntokun’s syndrome), but on further follow-up some have turned out to be variants of HSAN.

Reference

Cross References
Anaesthesia; Frontal lobe syndromes

**Anal Reflex**
Contraction of the external sphincter ani muscle in response to a scratch stimulus in the perianal region, testing the integrity of the S4/S5 roots, forms the anal or wink reflex. This reflex may be absent in some normal elderly individuals, and absence does not necessarily correlate with urinary incontinence. External anal responses to coughing and sniffing are part of a highly consistent and easily elicited polysynaptic reflex, whose characteristics resemble those of the conventional scratch-induced anal reflex.

Reference

Cross Reference
Reflexes

**Anarchic Hand**
- see ALIEN HAND, ALIEN LIMB

**Anarithmetia**
- see ACALCULIA

**Anarthria**
Anarthria is the complete inability to articulate words (cf. dysarthria). This is most commonly seen as a feature of the bulbar palsy of motor neurone disease.

A motor disorder of speech production with preserved comprehension of spoken and written language has been termed pure anarthria; this syndrome has also been labelled as aphemia, phonetic disintegration, apraxic dysarthria, cortical dysarthria, verbal apraxia, subcortical motor aphasia, pure motor aphasia, and small or mini-Broca’s aphasia. It reflects damage in the left frontal
Anismus

operculum, but with sparing of Broca’s area. A pure progressive anarthria or slowly progressive anarthria may result from focal degeneration affecting the frontal operculum bilaterally (so-called Foix–Chavany–Marie syndrome).

References

Cross References
Aphemia; Bulbar palsy; Dysarthria

Angioscotoma

Angioscotomata are shadow images of the superficial retinal vessels on the underlying retina, a physiological scotoma.

Cross Reference
Scotoma

Angor Animi

Angor animi is the sense of dying or the feeling of impending death. It may be experienced on awakening from sleep or as a somesthetic aura of migraine.

Reference

Cross Reference
Aura

Anhidrosis

Anhidrosis, or hypohidrosis, is a loss or lack of sweating. This may be due to primary autonomic failure or due to pathology within the posterior hypothalamus (‘sympathetic area’).

Anhidrosis may occur in various neurological disorders, including multiple system atrophy, Parkinson’s disease, multiple sclerosis, caudal to a spinal cord lesion, and in some hereditary sensory and autonomic neuropathies. Localized or generalized anhidrosis may be seen in Holmes–Adie syndrome, and unilateral anhidrosis may be seen in Horner’s syndrome if the symptomatic lesion is distal to the superior cervical ganglion.

Cross References
Holmes–Adie pupil, Holmes–Adie syndrome; Horner’s syndrome; Hyperhidrosis

Anismus

Anismus, also known as puborectalis syndrome, is paradoxical contraction of the external anal sphincter during attempted defaecation, leading to faecal retention and a complaint of constipation. This may occur as an idiopathic condition in isolation or as a feature of the off periods of idiopathic Parkinson’s disease. It
Anisocoria is thought to represent a focal dystonia and may be helped temporarily by local injections of botulinum toxin.

Reference

Cross References
Dystonia; Parkinsonism

Anisocoria
Anisocoria is an inequality of pupil size. This may be physiological (said to occur in up to 15% of the population), in which case the inequality is usually mild and does not vary with degree of ambient illumination; or pathological, with many possible causes.

- Structural:
  - Ocular infection, trauma, inflammation, surgery.

- Neurological:
  - Anisocoria greater in dim light or darkness suggests a sympathetic innervation defect (darkness stimulates dilatation of normal pupil). Affected pupil is constricted (miosis; oculosympathetic paresis), as in:
    - Horner’s syndrome;
    - Argyll Robertson pupil;
    - Cluster headache.
  - Anisocoria greater in bright light/less in dim light suggests a defect in parasympathetic innervation to the pupil. Affected pupil is dilated (mydriasis; oculoparasympathetic paresis), as in:
    - Holmes–Adie pupil (vermiform movements of the pupil margin may be visible with a slit lamp);
    - Oculomotor (III) nerve palsy (efferent path from Edinger–Westphal nucleus);
    - Mydriatic agents (phenylephrine, tropicamide);
    - Anticholinergic agents (e.g. asthma inhaler accidentally puffed into one eye).

Clinical characteristics and pharmacological testing may help to establish the underlying diagnosis in anisocoria.

Reference

Cross References
Argyll Robertson pupil; Holmes–Adie pupil, Holmes–Adie syndrome; Horner’s syndrome; Miosis; Mydriasis

Annular Scotoma
An annular or ring scotoma suggests retinal disease, as in retinitis pigmentosa or cancer-associated retinopathy (paraneoplastic retinal degeneration).
Anosmia

Anosmia is the inability to perceive smells due to damage to the olfactory pathways (olfactory neuroepithelium, olfactory nerves, rhinencephalon). Olfaction may be tested with kits containing specific odours (e.g. clove, turpentine); each nostril should be separately tested. Unilateral anosmia may be due to pressure on the olfactory bulb or tract, e.g. due to a subfrontal meningioma.

Anosmia may be congenital (e.g. Kallman’s syndrome, hypogonadotrophic hypogonadism, a disorder of neuronal migration) or, much more commonly, acquired. Rhinological disease (allergic rhinitis, coryza) is by far the most common cause; this may also account for the impaired sense of smell in smokers. Head trauma is the most common neurological cause, due to shearing off of the olfactory fibres as they pass through the cribriform plate. Recovery is possible in this situation due to the capacity for neuronal and axonal regeneration within the olfactory pathways. Olfactory dysfunction is also described in Alzheimer’s disease and Parkinson’s disease, possibly as an early phenomenon, due to pathological involvement of olfactory pathways. Patients with depression may also complain of impaired sense of smell. Loss of olfactory acuity may be a feature of normal ageing.

Cross References
Retinopathy; Scotoma; Visual field defects

Anomia

Anomia or dysnomia is a deficit in naming or word-finding. This may be detected as abrupt cut-offs in spontaneous speech with circumlocutions and/or paraphasic substitutions. Formal tests of naming are also available (e.g. Graded Naming Test). Patients may be able to point to named objects despite being unable to name them, suggesting a problem in word retrieval but with preserved comprehension. They may also be able to say something about the objects they cannot name (e.g. ‘flies in the sky’ for kite), suggesting preserved access to the semantic system.

Category-specific anomias have been described, e.g. for colour (cf. achromatopsia).

Anomia occurs with pathologies affecting the left temporoparietal area, but since it occurs in all varieties of aphasia is of little precise localizing or diagnostic value. The term anomic aphasia is reserved for unusual cases in which a naming problem overshadows all other deficits. Anomia may often be seen as a residual deficit following recovery from other types of aphasia. Anomia may occur with any dominant hemisphere space-occupying lesion, and as a feature of semantic dementia, being more prominent in this condition than in Alzheimer’s disease.

References

Cross References
Aphasia; Circumlocution; Paraphasia
Anosodiaphoria

Babinski (1914) used the term anosodiaphoria to describe a disorder of body schema in which patients verbally acknowledge a clinical problem (e.g. hemiparesis) but fail to be concerned by it. Anosodiaphoria usually follows a stage of anosognosia.

_La belle indifférence_ describes a similar lack of concern for acknowledged disabilities which are psychogenic.

**References**


**Cross References**

Anosognosia; _Belle indifférence_; Personification of paralyzed limbs

**Anosognosia**

Anosognosia refers to a patient’s unawareness or denial of their illness. The term was first used by von Monakow (1885) and has been used to describe denial of blindness (Anton’s syndrome), deafness, hemiplegia (Babinski), hemianopia, aphasia, and amnesia. Some authorities would question whether this unawareness is a true agnosia or rather a defect of higher-level cognitive integration (i.e. perception).

Anosognosia with hemiplegia most commonly follows right hemisphere injury (parietal and temporal lobes) and may be associated with left hemineglect and left-sided hemianopia; it is also described with right thalamic and basal ganglia lesions. Many patients with posterior aphasia (Wernicke type) are unaware that their output is incomprehensible or jargon, possibly through a failure to monitor their own output. Cerebrovascular disease is the most common pathology associated with anosognosia, although it may also occur with neurodegenerative disease, for example, the cognitive anosognosia in some patients with Alzheimer’s disease.

The neuropsychological mechanisms of anosognosia are unclear: the hypothesis that it might be accounted for by personal neglect (asomatognosia), which is also more frequently observed after right hemisphere lesions, would seem to have been disproved experimentally by studies using selective hemisphere anaesthesia in which the two may be dissociated, a dissociation which may also be observed clinically. In Alzheimer’s disease, anosognosia may be related to memory dysfunction and executive dysfunction.

At a practical level, anosognosia may lead to profound difficulties with neurorehabilitation. Temporary resolution of anosognosia has been reported following vestibular stimulation (e.g. with caloric testing).
Anteflexion

References

Cross References
Agnosia; Anosodiaphoria; Asomatognosia; Cortical blindness; Extinction; Jargon aphasia; Misoplegia; Neglect; Personification of paralyzed limbs; Somatoparaphrenia

Anserina
Autonomically mediated piloerection and thermoconstriction may produce ‘goosebumps’, cold and bumpy skin which may be likened to that of a plucked goose. Loss of anserina may be a feature of some autonomic disorders.

Antecollis
Antecollis (anterocollis) is forward flexion of the neck. It may be a feature of multiple system atrophy (cf. retrocollis in progressive supranuclear palsy), a sustained dystonic posture in advanced Parkinson’s disease, and, unusually, in spasmodic torticollis.

Forward flexion of the head onto the chest is a feature in the ‘dropped head syndrome’.

Reference

Cross References
Dropped head syndrome; Retrocollis; Torticollis

Anteflexion
Anteflexion is forward flexion of the trunk, as typical of the stooped posture seen in Parkinson’s disease.

Cross Reference
Parkinsonism
**Anton’s Syndrome**

Anton’s syndrome is cortical blindness accompanied by denial of the visual defect (visual anosognosia), with or without confabulation. The syndrome most usually results from bilateral posterior cerebral artery territory lesions causing occipital or occipitoparietal infarctions but has occasionally been described with anterior visual pathway lesions associated with frontal lobe lesions. It may also occur in the context of dementing disorders or delirium.

**References**


**Cross References**

Agnosia, Anosognosia, Confabulation, Cortical blindness

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**Anwesenheit**

A vivid sensation of the presence of somebody either somewhere in the room or behind the patient has been labelled as *anwesenheit* (German: presence), presence hallucination, minor hallucination, or extracampine hallucination. This phenomenon is relatively common in Parkinson’s disease, occurring in isolation or associated with formed visual hallucinations.

**References**

Chan D, Rossor MN. “- but who is that on the other side of you?” Extracampine hallucinations revisited. *Lancet* 2002; **360**: 2064–2066.


**Cross References**

Hallucination; Parkinsonism

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**Apallic Syndrome**

- see VEGETATIVE STATES

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**Apathy**

Apathy is a neurobehavioural disorder which may be characterized by:

1. A lack of motivation relative to the patient’s previous level of functioning or the standards of age and culture.
2. Presence of at least one of the following:
   a. Diminished goal-directed behaviour: lack of effort, dependency on others to structure activity;
   b. Diminished goal-directed cognition: lack of interest, concern about personal problems;
   c. Diminished concomitants of goal-directed behaviour: unchanging affect, lack of emotional responsiveness.
An Apathy Inventory has been developed based on these criteria. Hence, listlessness, paucity of spontaneous movement (akinesia) or speech (mutism), and lack of initiative, spontaneity, and drive may be features of apathy. These are also all features of the abulic state, and it has been suggested that apathy and abulia represent different points on a continuum of motivational and emotional deficit, abulia being at the more severe end. The diminished motivation of apathy should not be attributable to impaired level of consciousness, emotional distress, or cognitive impairment although it may coexist with the latter, as in Alzheimer’s disease. Apathy is a specific neuropsychiatric syndrome, distinct from depression.

Apathy may be observed in diseases affecting frontal–subcortical structures, for example, in the frontal lobe syndrome affecting the frontal convexity, or following multiple vascular insults to paramedian diencephalic structures (thalamus, subthalamus, posterior lateral hypothalamus, mesencephalon) or the posterior limb of the internal capsule; there may be associated cognitive impairment of the so-called subcortical type in these situations (e.g. in Huntington’s disease). Apathy is also described following amphetamine or cocaine withdrawal, in neuroleptic-induced akinesia and in psychotic depression. Selective serotonin-reuptake inhibitors may sometimes be helpful in the treatment of apathy.

References

Cross References
Abulia; Akinetic mutism; Dementia; Frontal lobe syndromes

Aphasia
Aphasia, or dysphasia, is an acquired loss or impairment of language function. Language may be defined as the complex system of symbols used for communication (including reading and writing), encompassing various linguistic components (phonetic, phonemic, semantic/lexical, syntactic, pragmatic), all of which are dependent on dominant hemisphere integrity. Non-linguistic components of language (emotion, inflection, cadence), collectively known as prosody, may require contributions from both hemispheres. Language is distinguished from speech (oral communication), disorders of which are termed dysarthria or anarthria. Dysarthria and aphasia may coexist but are usually separable.

Clinical assessment of aphasia requires analysis of the following features, through listening to the patient’s spontaneous speech, asking questions or giving commands, and asking the patient to repeat, name, read and write:
Aphasia

- **Fluency**: is output effortful, laboured, with agrammatism and dysprosody (non-fluent); or flowing, with paraphasias and neologisms (fluent)?
- **Comprehension**: spared or impaired?
- **Repetition**: preserved or impaired?
- **Naming**: preserved or impaired?
- **Reading**: evidence of alexia?
- **Writing**: evidence of agraphia?

These features allow definition of various types of aphasia (see table and specific entries; although it should be noted that some distinguished neurologists have taken the view that no satisfactory classification of the aphasias exists (Critchley)). For example, motor (‘expressive’) aphasias are characterized by non-fluent verbal output, with intact or largely unimpaired comprehension, whereas sensory (‘receptive’) aphasias demonstrate fluent verbal output, often with paraphasias, sometimes jargon, with impaired comprehension. Conduction aphasia is marked by relatively normal spontaneous speech (perhaps with some paraphasic errors), but a profound deficit of repetition. In transcortical motor aphasia spontaneous output is impaired but repetition is intact.

<table>
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<th>Wernicke</th>
<th>Conduction</th>
<th>Transcortical: motor/sensory</th>
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N = normal; ↓ = impaired

Aphasias most commonly follow a cerebrovascular event: the specific type of aphasia may change with time following the event, and discrepancies may be observed between classically defined clinicoanatomical syndromes and the findings of everyday practice. Aphasia may also occur with space-occupying lesions and in neurodegenerative disorders, often with other cognitive impairments (e.g. Alzheimer’s disease) but sometimes in isolation (primary non-fluent aphasia, semantic dementia).

**References**


Aphonia


Cross References
Agrammatism; Agraphia; Alexia; Anomia; Aprosodia, Aprosody; Broca’s aphasia; Circumlocution; Conduction aphasia; Conduit d’approche; Crossed aphasia; Dysarthria; Jargon aphasia; Neologism; Optic aphasia; Paraphasia; Transcortical aphasias; Wernicke’s aphasia

Aphemia
Aphemia was the name originally given by Broca to the language disorder subsequently named ‘Broca’s aphasia’. The term is now used to describe a motor disorder of speech production with preserved comprehension of spoken and written language. This syndrome has also been called phonetic disintegration (cf. phonemic disintegration), pure anarthria, apraxic dysarthria, cortical dysarthria, verbal apraxia, subcortical motor aphasia, alalia, pure motor aphasia, small or mini-Broca’s aphasia, and kinetic speech production disorder, reflecting the differing views as to the nature of the underlying disorder (aphasia, dysarthria, apraxia). Aphemia probably encompasses at least some cases of the ‘foreign accent syndrome’, in which altered speech production and/or prosody makes speech output sound foreign. Such conditions may stand between pure disorders of speech (i.e. dysarthrias) and of language (i.e. aphasias). They usually reflect damage in the left frontal operculum, but sparing Broca’s area.

References


Cross References
Anarthria; Aphasia; Aprosodia, Aprosody; Dysarthria; Phonemic disintegration; Speech apraxia

Aphonia
Aphonia is loss of the sound of the voice, necessitating mouthing or whispering of words. As for dysphonia, this most frequently follows laryngeal inflammation, although it may follow bilateral recurrent laryngeal nerve palsy. Dystonia of the abductor muscles of the larynx can result in aphonie segments of speech (spasmodic aphony or abductor laryngeal dystonia); this may be diagnosed by
hearing the voice fade away to nothing when asking the patient to keep talking; patients may comment that they cannot hold any prolonged conversation. Aphonia of functional or hysterical origin is also recognized.

Aphonia should be differentiated from mutism, in which patients make no effort to speak, and anarthria in which there is a failure of articulation.

**Cross References**
Anarthria; Dysphonia; Mutism

**Applause Sign**
To elicit the applause sign, also known as the clapping test or three clap test, the patient is asked to clap the hands three times. The tendency to clap more than three times, even when demonstrated by the examiner, is said to be specific to striatal dysfunction and is seen in progressive supranuclear palsy to a greater extent than in Parkinson’s disease, but not in frontotemporal dementia.

**References**

**Aposiopesis**
Critchely used this term to denote a sentence which is started but not finished, as in the aphasia associated with dementia.

**Reference**

**Cross Reference**
Aphasia

**Apraxia**
Apraxia or dyspraxia is a disorder of movement characterized by the inability to perform a voluntary motor act despite an intact motor system (i.e. no ataxia, weakness) and without impairment in level of consciousness. Automatic/reflex actions are preserved, hence there is a voluntary–automatic dissociation; some authors see this as critical to the definition of apraxia. Different types of apraxia have been delineated, the standard classification being that of Liepmann (1900):

- **Ideational apraxia, conceptual apraxia:**
  A deficit in the conception of a movement; this frequently interferes with daily motor activities and is not facilitated by the use of objects; there is often an associated aphasia.

- **Ideomotor apraxia (IMA):**
  A disturbance in the selection of elements that constitute a movement (e.g. pantomiming the use of tools); in contrast to ideational apraxia, this is a ‘clinical’ disorder inasmuch as it does not greatly interfere with everyday activities; moreover, use of objects may facilitate movement;
 Apraxia  

it may often be manifest as the phenomenon of using body part as object, e.g. in demonstrating how to use a toothbrush or how to hammer a nail, a body part is used to represent the object (finger used as toothbrush, fist as hammer).

- **Limb-kinetic, or melokinetic, apraxia:**
  
  Slowness, clumsiness, awkwardness in using a limb, with a temporal decomposition of movement; difficult to disentangle from pure motor deficits associated with corticospinal tract lesions.

Apraxia may also be defined anatomically:

- **Parietal (posterior):**
  
  Ideational and ideomotor apraxia are seen with unilateral lesions of the inferior parietal lobule (most usually of the left hemisphere) or premotor area of the frontal lobe (Brodmann areas 6 and 8).

- **Frontal (anterior):**
  
  Unilateral lesions of the supplementary motor area are associated with impairment in tasks requiring bimanual coordination, leading to difficulties with alternating hand movements, drawing alternating patterns (e.g. m n m n in joined up writing: alternating sequences test, Luria figures). This may be associated with the presence of a grasp reflex and alien limb phenomena (limb-kinetic type of apraxia).

Apraxia is more common and severe with left hemisphere lesions.

Difficulties with the clinical definition of apraxia persist, as for the agnosias. For example, ‘dressing apraxia’ and ‘constructional apraxia’ are now considered visuospatial problems rather than true apraxias. Likewise, some cases labelled as eyelid apraxia or gait apraxia are not true ideational apraxias. The exact nosological status of speech apraxia also remains tendentious.

**References**


Liepmann H. *Das Krankheitsbild der Apraxie (“motorischen Asymbolie”).* Berlin: Karger, 1900.

**Cross References**

Alien hand, Alien limb; Body part as object; Crossed apraxia; Dysdiadochokinesia; Eyelid apraxia; Forced groping; Frontal lobe syndromes; Gait apraxia; Grasp reflex; Optic ataxia; Speech apraxia
Aprosexia
Aprosexia is a syndrome of psychomotor inefficiency, characterized by complaints of easy forgetting, for example, of conversations as soon as they are finished, material just read, or instructions just given. There is difficulty keeping the mind on a specific task, which is forgotten if the patient happens to be distracted by another task. These difficulties, into which the patient has insight and often bitterly complains of, are commonly encountered in the memory clinic. They probably represent a disturbance of attention or concentration, rather than being a harbinger of dementia. These patients generally achieve normal scores on formal psychometric tests (and indeed may complain that these assessments do not test the function they are having difficulty with). Concurrent sleep disturbance, irritability, and low mood are common and may reflect an underlying affective disorder (anxiety, depression) which may merit specific treatment.

Cross References
Attention; Dementia

Aprosodia, Aprosody
Aprosodia or aprosody (dysprosodia, dysprosody) is a defect in or absence of the ability to produce or comprehend speech melody, intonation, cadence, rhythm, and accentuations, in other words the non-linguistic aspects of language which convey or imply emotion and attitude. Aprosodia may be classified, in a manner analogous to the aphasias, as:

- **Sensory (posterior):** Impaired comprehension of the emotional overtones of spoken language or emotional gesturing, also known as affective agnosia; this may be associated with visual extinction and anosognosia, reflecting right posterior temporoparietal region pathology.

- **Expressive/motor (anterior):** An inability to produce emotional overtones (‘emotional dysprosody’, sometimes confusingly referred to as speech dyspraxia); this may occur in isolation with right-sided anterior lesions or in association with linguistic aspects of aphasia such as agrammatism with anterior left hemisphere damage.

References

Cross References
Agnosia: Agrammatism; Anosognosia; Aphasia; Aphemia; Broca’s aphasia; Fisher’s sign; Visual extinction

Arc de Cercle
- see OPISTHOTONOS
**Arcuate Scotoma**
An arcuate scotoma suggests retinal or optic nerve disease, such as glaucoma, acute ischaemic optic neuropathy, or the presence of drusen.

**Cross References**
Retinopathy; Scotoma

**Areflexia**
Areflexia is an absence or a loss of tendon reflexes. This may be physiological, in that some individuals never demonstrate tendon reflexes; or pathological, reflecting an anatomical interruption or physiological dysfunction at any point along the monosynaptic reflex pathway which is the neuroanatomical substrate of phasic stretch reflexes. Sudden tendon stretch, as produced by a sharp blow from a tendon hammer, activates muscle spindle Ia afferents which pass to the ventral horn of the spinal cord, there activating α-motor neurones, the efferent limb of the reflex, so completing the monosynaptic arc. Hence, although reflexes are typically regarded as part of the examination of the motor system, reflex loss may also occur in ‘sensory’ disorders, affecting the Ia afferents from the muscle spindle. It is often possible to ‘hear’ that reflexes are absent from the thud of tendon hammer on tendon.

Areflexia is most often encountered in disorders of lower motor neurones, specifically radiculopathies, plexopathies, and neuropathies (axonal and demyelinating). Areflexia may also occur in neuromuscular junction disorders, such as the Lambert–Eaton myasthenic syndrome, in which condition the reflexes may be ‘restored’ following forced muscular contraction (facilitation). Transient areflexia may be seen in central nervous system disorders, such as cataplexy, and in acute spinal cord syndromes (‘spinal shock’, e.g. acute compression, acute inflammatory myelopathy).

**Cross References**
Cataplexy; Facilitation; Hyporeflexia; Lower motor neurone (LMN) syndrome; Plexopathy; Radiculopathy; Reflexes

**Argyll Robertson Pupil (ARP)**
The Argyll Robertson pupil is small (miosis) and irregular. It fails to react to light (reflex iridoplegia), but does constrict to accommodation (when the eyes converge). In other words, there is light-near pupillary dissociation (ARP = accommodation reaction preserved). Since the light reflex is lost, testing for the accommodation reaction may be performed with the pupil directly illuminated: this can make it easier to see the response to accommodation, which is often difficult to observe when the pupil is small or in individuals with a dark iris. There may be an incomplete response to mydriatic drugs. Although pupil involvement is usually bilateral, it is often asymmetric, causing anisocoria.

The Argyll Robertson pupil was originally described in the context of neurosyphilis, especially tabes dorsalis. If this pathological diagnosis is suspected, a helpful clinical concomitant is the associated loss of deep pain sensation, as assessed, for example, by vigorously squeezing the Achilles tendon (Abadie’s sign). There are, however, a number of recognized causes of ARP besides neurosyphilis, including
- Multiple sclerosis;
- Encephalitis;
- Diabetes mellitus;
- Syringobulbia;
- Sarcoidosis;
- Lyme disease;
- Pinealoma;
- Herpes zoster;
- Hereditary motor and sensory neuropathies (Charcot–Marie Tooth disease; Dejerine–Sottas hypertrophic neuropathy).

Miosis and pupil irregularity are inconstant findings in some of these situations, in which case the term ‘pseudo-Argyll Robertson pupil’ may be preferred. The neuroanatomical substrate of the Argyll Robertson pupil is uncertain. A lesion in the tectum of the (rostral) midbrain proximal to the oculomotor nuclei has been suggested. In multiple sclerosis and sarcoidosis, magnetic resonance imaging has shown lesions in the periaqueductal grey matter at the level of the Edinger–Westphal nucleus, but these cases lacked miosis and may therefore be classified as pseudo-Argyll Robertson pupil. Some authorities think that a partial oculomotor (III) nerve palsy or a lesion of the ciliary ganglion is a more likely cause.

References

Cross References
Abadie’s sign; Anisocoria; Light-near pupillary dissociation; Miosis; Pseudo-Argyll Robertson pupil

Arm Drop
‘Arm drop’, or the ‘face–hand test’, has been suggested as a useful diagnostic test if hemiparesis or upper limb monoparesis is suspected to be psychogenic: the examiner lifts the paretic hand directly over the patient’s face and drops it. It is said that in organic weakness the hand will hit the face, whereas patients with functional weakness avoid this consequence. However, the validity and reliability of this ‘avoidance testing manoeuvre’ has never been examined; its clinical value is therefore doubtful.

Reference

Cross References
Babinski’s trunk–thigh test; Functional weakness and sensory disturbance; Hoover’s sign
Asemia
Asemia is an inability to indicate by signs or spoken language. The term was invented in the nineteenth century (Hamilton) as an alternative to aphasia, since in many cases of the latter there is more than a loss of speech, including impaired pantomime (apraxia) and in symbolizing the relationships of things. Hughlings Jackson approved of the term but feared it was too late to displace the word aphasia.

Reference

Cross References
Aphasia, Apraxia

Asomatognosia
Asomatognosia is a lack of regard for a part, or parts, of the body, most typically failure to acknowledge the existence of a hemiplegic left arm. Asomatognosia may be verbal (denial of limb ownership) or non-verbal (failure to dress or wash limb). All patients with asomatognosia have hemispatial neglect (usually left), hence this would seem to be a precondition for the development of asomatognosia; indeed, for some authorities asomatognosia is synonymous with personal neglect. Attribution of the neglected limb to another person is known as somatoparaphrenia.

The neuroanatomical correlate of asomatognosia is damage to the right supramarginal gyrus and posterior corona radiata, most commonly due to a cerebrovascular event. Cases with right thalamic lesions have also been reported. The predilection of asomatognosia for the left side of the body may simply be a reflection of the aphasic problems associated with left-sided lesions that might be expected to produce asomatognosia for the right side.

Asomatognosia is related to anosognosia (unawareness or denial of illness) but the two are dissociable on clinical and experimental grounds. Some authorities consider asomatognosia as a form of confabulation.

Reference

Cross References
Anosognosia; Confabulation; Neglect; Somatoparaphrenia

Astasia
- see CATAPLEXY
Astasia–Abasia

Astasia–abasia is the name which has sometimes been given to a disorder of gait characterized by impaired balance (disequilibrium), wide base, shortened stride, start/turn hesitation, and freezing. The term has no standardized definition and hence may mean different things to different observers; it has also been used to describe a disorder characterized by inability to stand or walk despite normal leg strength when lying or sitting, believed to be psychogenic (although gait apraxia may have similar features). Modern clinical classifications of gait disorders subsume astasia–abasia under the categories of subcortical disequilibrium and frontal disequilibrium, i.e. gait disorders with prominent disequilibrium or impaired postural control. A transient inability to sit or stand despite normal limb strength may be seen after an acute thalamic lesion (thalamic astasia).

References

Cross Reference
Gait apraxia

Astereognosis

Astereognosis is the failure to recognize a familiar object, such as a key or a coin, palpated in the hand with the eyes closed, despite intact primary sensory modalities. Description of qualities such as the size, shape, and texture of the object may be possible. Hence, this is a failure of higher-order (i.e. cortical) processing and is associated with lesions of the posterior parietal lobe (postcentral gyrus) association cortex. There may be associated impairments of two-point discrimination and graphaesthesia (cortical sensory syndrome). Astereognosis was said to be invariably present in the original description of the thalamic syndrome by Dejerine and Roussy.

Some authorities recommend the terms stereoanaesthesia or stereohypaesthesia as more appropriate descriptors of this phenomenon, to emphasize that this may be a disorder of perception rather than a true agnosia (for a similar debate in the visual domain, see Dysexopsia).

Cross References
Agnosia; Dysexopsia; Graphaesthesia; Two-point discrimination

Asterixis

Asterixis is a sudden, brief, arrhythmic lapse of sustained posture due to involuntary interruption in muscle contraction. It is most easily demonstrated by observing the dorsiflexed hands with arms outstretched (i.e. the motion to indicate ‘stop’), lapses being seen as flicking or flapping movements of the hands (‘flapping tremor’). Movement is associated with EMG silence in antigravity muscles for 35–200 ms. These features distinguish asterixis from tremor and myoclonus; the phenomenon has previously been described as negative myoclonus or negative tremor. Asterixis may be bilateral or unilateral. Recognized causes of asterixis include
- Hepatic encephalopathy (‘liver flap’);
- Hypercapnia;
- Uraemia;
- Drug-induced, e.g. anticonvulsants, levodopa;
- Structural brain lesions: thalamic lesions (haemorrhage, thalamotomy).

Unilateral asterixis has been described in the context of stroke, contralateral to lesions of the midbrain (involving corticospinal fibres, medial lemniscus), thalamus (ventroposterolateral nucleus), primary motor cortex, and parietal lobe; and ipsilateral to lesions of the pons or medulla.

References

Cross References
Encephalopathy; Myoclonus; Tremor

Asthenopia
Asthenopia, literally ‘weak vision’, is frequently used to describe ‘eye strain’ due to uncorrected or incorrectly corrected refractive errors, such as hyperopia (far-sightedness) or overcorrected myopia. Such refractive errors are sometimes blamed for headache.

Asynergy
Asynergia or dyssynergia is lack or impairment of synergy of sequential muscular contraction in the performance of complex movements, such that they seem to become broken up into their constituent parts, so-called decomposition of movement. This may be evident when performing rapid alternating hand movements. Dyssynergy of speech may also occur, a phenomenon sometimes termed scanning speech or scanning dysarthria. This is typically seen in cerebellar syndromes, most often those affecting the cerebellar hemispheres, and may coexist with other signs of cerebellar disease such as ataxia, dysmetria, and dysdiadochokinesia.

Cross References
Ataxia; Cerebellar syndromes; Dysarthria; Dysdiadochokinesia; Dysmetria; Scanning speech

Ataxia
Ataxia or dystaxia refers to a lack of coordination of voluntary motor acts, impairing their smooth performance. The rate, range, timing, direction, and force of movement may be affected. Ataxia is used most frequently to refer to a cerebellar problem, but sensory ataxia, optic ataxia, and frontal ataxia are also described, so it is probably best to qualify ataxia rather than to use the word in isolation.
A

Ataxia

- **Cerebellar ataxia:**
  Defective timing of agonist and antagonist muscle contraction (asynergia) produces jerking, staggering, inaccurate movements (decomposition of movement), which may manifest as intention tremor, dysmetria (past pointing), dysdiadochokinesia, ataxic dysarthria (sometimes known as scanning speech, although this also has other connotations), excessive rebound phenomenon, macrographia, head tremor (titubation), gait ataxia, and abnormal eye movements (nystagmus, square wave jerks, saccadic intrusions). There may be concurrent limb hypotonia. Cerebellar hemisphere lesions cause ipsilateral limb ataxia (hemiataxia; ataxia on finger-to-nose, finger chase, and/or heel–shin testing) whereas midline cerebellar lesions involving the vermis produce selective truncal and gait ataxia. An International Cooperative Ataxia Rating Scale has been developed to assess the efficacy of treatments for cerebellar ataxia.

- **Sensory ataxia:**
  Results from impaired proprioception and may be seen in disease of the dorsal (posterior) columns of the spinal cord (hence ‘spinal ataxia’), sensory neuropathies, and neuronopathies affecting the dorsal root ganglia. It is markedly exacerbated by removal of visual cues (e.g. as in Romberg’s sign), unlike the situation with cerebellar ataxia, and may also lead to pseudoathetosis.

- **Optic ataxia:**
  Misreaching for visually presented targets, with dysmetria, due to a parieto-occipital lesion, as seen in Balint’s syndrome.

- **‘Frontal ataxia’:**
  Similar to, and sometimes indistinguishable from, cerebellar ataxia, but results from lesions of the contralateral frontal cortex or frontopontine fibres, often from tumours invading the frontal lobe or corpus callosum. These fibres run in the corticopontocerebellar tract, synapsing in the pons before passing through the middle cerebellar peduncle to the contralateral cerebellar hemisphere.

Triple ataxia, the rare concurrence of cerebellar, sensory, and optic types of ataxia, may be associated with an alien limb phenomenon (sensory type).

There are many causes of cerebellar ataxia, including

- **Inherited:**
  *Autosomal recessive:* Friedreich’s ataxia, ataxia with isolated vitamin E deficiency, ataxia with oculomotor apraxia (types 1 and 2);
  *Autosomal dominant:* clinically ADCA types I, II, and III, now reclassified genetically as spinocerebellar ataxias: types 1–31 now described;
  *Episodic ataxias:* channelopathies involving potassium (type 1) and calcium (type 2) ion channels;
  Mitochondrial disorders;
  Huntington’s disease;
  Dentatorubropallidoluysian atrophy (DRPLA);
Inherited prion diseases, especially Gerstmann–Straussler–Scheinker (GSS) syndrome.

- **Acquired:**

  *Cerebrovascular events (infarct, haemorrhage):* usually cause hemiataxia; postanoxic cerebellar ataxia;  
  *Inflammatory: demyelination:* multiple sclerosis, Miller Fisher variant of Guillain–Barré syndrome, central pontine and extrapontine myelinolysis;  
  *Inflammatory: infection:* cerebellitis with Epstein–Barr virus; encephalitis with *Mycoplasma*; HIV;  
  *Neoplasia:* tumours, paraneoplastic syndromes;  
  *Neurodegeneration:* one variant of multiple system atrophy (MSA-C); prion diseases (Brownell–Oppenheimer variant of sporadic Creutzfeldt–Jakob disease, kuru); idiopathic late-onset cerebellar ataxia;  
  *Drugs/toxins,* e.g. alcohol, phenytoin.  
  *Metabolic:* vitamin E deficiency, thiamine deficiency (Wernicke’s encephalopathy), gluten ataxia, hypothyroidism (debatable).

**References**


**Cross References**  
Alien hand, Alien limb; Asynergia; Balint’s syndrome; Cerebellar syndromes; Dysarthria; Dysdiadochokinesia; Dysmetria; Head tremor; Hemiataxia; Hypotonia, Hypotonus; Macrographia; Nystagmus; Optic ataxia; Proprioception; Pseudoathetosis; Rebound phenomenon; Rombergism, Romberg's sign; Saccadic intrusion, Saccadic pursuit; Scanning speech; Square wave jerks; Tandem walking; Tremor

**Ataxic Hemiparesis**

Ataxic hemiparesis is a syndrome of ipsilateral hemiataxia and hemiparesis, the latter affecting the leg more severely than the arm (crural paresis). There may be additional dysarthria, nystagmus, paraesthesia, and pain.

This syndrome is caused by lacunar (small deep) infarcts in the contralateral basis pons at the junction of the upper third and lower two-thirds. It may also be seen with infarcts in the contralateral thalamocapsular region, posterior limb of the internal capsule (anterior choroidal artery syndrome), red nucleus, and the paracentral region (anterior cerebral artery territory). Sensory loss is an indicator of capsular involvement; pain in the absence of other sensory features is an indicator of thalamic involvement.
Ataxic Nystagmus

References

Cross References
Ataxia; Hemiataxia; Hemiparesis; Pseudochoreoathetosis

Ataxic Nystagmus
- see INTERNUCLEAR OPHTHALMOLEGIA; NYSTAGMUS

Athetosis
Athetosis is the name sometimes given to an involuntary movement disorder characterized by slow, sinuous, purposeless, writhing movements, often more evident in the distal part of the limbs. Athetosis often coexists with the more flowing, dance-like movements of chorea, in which case the movement disorder may be described as choreoathetosis. Indeed the term athetosis is now little used except in the context of ‘athetoid cerebral palsy’. Athetoid-like movements of the outstretched hands may also been seen in the presence of sensory ataxia (impaired proprioception) and are known as pseudoathetosis or pseudochoreoathetosis. Choreoathetoid movements result from disorders of the basal ganglia.

References

Cross References
Chorea, Choreoathetosis; Pseudoathetosis; Pseudochoreoathetosis

Athymhormia
Athym(h)ormia, also known as the robot syndrome, is a name given to a form of abulia or akinetic mutism in which there is loss of self-autoactivation. Clinically there is a marked discrepancy between heteroactivation, behaviour under the influence of exogenous stimulation, which is normal or almost normal, and autoactivation. Left alone, patients are akinetic and mute, a state also known as loss of psychic self-activation or pure psychic akinesia. It is associated with bilateral deep lesions of the frontal white matter or of the basal ganglia, especially the globus pallidus. Athymhormia is thus environment-dependent, patients normalizing initiation and cognition when stimulated, an important differentiation from apathy and akinetic mutism.

Reference
**Cross References**

Abulia; Akinetic mutism; Apathy

**Atrophy**

Atrophy is a wasting or thinning of tissues. The term is often applied to wasted muscles, usually in the context of lower motor neurone pathology (in which case it may be synonymous with amyotrophy), but also with disuse. Atrophy develops more quickly after lower, as opposed to upper, motor neurone lesions. It may also be applied to other tissues, such as subcutaneous tissue (as in hemifacial atrophy). Atrophy may sometimes be remote from the affected part of the neuraxis, hence a false-localizing sign, for example, wasting of intrinsic hand muscles with foramen magnum lesions.

**Cross References**

Amyotrophy; ‘False-localizing signs’; Hemifacial atrophy; Lower motor neurone (LMN) syndrome; Wasting

**‘Attended Alone’ Sign**

Collateral history is crucial in assessing cognitive disorders, especially complaints of memory impairment, for which reason individuals referred to memory clinics are usually asked to bring with them a spouse, relative, or friend who knows them well to provide such history. Failure to attend with an informant, the ‘attended alone’ sign, is a robust (i.e. very sensitive, > 0.95) marker of the absence of dementia.

**Reference**


**Cross Reference**

Dementia

**Attention**

Attention is a distributed cognitive function, important for the operation of many other cognitive domains; the terms concentration, vigilance, and persistence may be used synonymously with attention. Distinction may be made between different types of attention, as follows:

- Sustained;
- Selective;
- Divided/executive function.

It is generally accepted that attention is effortful, selective, and closely linked to intention.

Impairment of attentional mechanisms may lead to distractability (with a resulting complaint of poor memory, perhaps better termed aprosexia), disorientation in time and place, perceptual problems, and behavioural problems (e.g. disinhibition), as in the cardinal disorder of attention, delirium.

The neuroanatomical substrates of attention encompass the ascending reticular activating system of the brainstem, the thalamus, and the prefrontal (multimodal association) cerebral cortex (especially on the right). Damage to any of these areas may cause impaired attention.
Auditory Agnosia

Attentional mechanisms may be tested in a variety of ways. Those adapted to ‘bedside’ use all essentially look for a defect in selective attention, also known as working memory or short-term memory (although this does not necessarily equate with lay use of the term ‘short-term memory’):

- Orientation in time/place;
- Digit span forwards/backwards;
- Reciting months of the year backwards, counting back from 30 to 1;
- Serial sevens (serial subtraction of 7 from 100, = 93, 86, 79, 72, 65).

In the presence of severe attentional disorder (as in delirium) it is difficult to make any meaningful assessment of other cognitive domains (e.g. memory).

Besides delirium, attentional impairments may be seen following head injury, and in ostensibly ‘alert’ patients, e.g. with Alzheimer’s disease (the dysexecutive syndrome of impaired divided attention).

References

Cross References
Aprosexia; Delirium; Dementia; Disinhibition; Dysexecutive syndrome; Frontal lobe syndromes; Pseudodementia

Auditory Agnosia

Auditory agnosia refers to an inability to appreciate the meaning of sounds despite normal perception of pure tones as assessed by audiological examination. This agnosia may be for either verbal material (pure word deafness) or non-verbal material, either sounds (bells, whistles, animal noises) or music (amusia, of receptive or sensory type).

Cross References
Agnosia; Amusia; Phonagnosia; Pure word deafness

Auditory–Visual Synaesthesia

This name has been given to the phenomenon of sudden sound-evoked light flashes in patients with optic nerve disorders. This may be equivalent to noise-induced visual phosphenes or sound-induced photisms. It is not certain that this phenomenon meets suggested criteria for synaesthesia.

Reference

Cross References
Phosphene; Synaesthesia
Augmentation

The term augmentation may be used to describe a phenomenon seen in Lambert–Eaton myasthenic syndrome (LEMS), namely, an increase in strength of affected muscles detected in the first few seconds of maximal voluntary contraction, one feature, along with facilitation, of posttetanic potentiation. This may also be known as Lambert’s sign.

Augmentation also refers to the paradoxical worsening of the symptoms of restless legs syndrome with dopaminergic treatment, manifesting with earlier onset of symptoms in the evening or afternoon, shorter periods of rest to provoke symptoms, greater intensity of symptoms when they occur, spread of symptoms to other body parts such as the arms, and decreased duration of benefit from medication.

Cross Reference
Facilitation

Aura

An aura is a brief feeling or sensation, lasting seconds to minutes, occurring immediately before the onset of a paroxysmal neurological event such as an epileptic seizure or a migraine attack (migraine with aura, ‘classical migraine’), ‘warning’ of its imminent presentation, although auras may also occur in isolation. An aura indicates the focal onset of neurological dysfunction. Auras are exclusively subjective, and may be entirely sensory, such as the fortification spectra (teichopsia) of migraine, or more complex, labelled psychosensory or experiential, as in certain seizures. Epileptic auras may be classified into subgroups:

- **Somatosensory:**
  - for example, paraesthesia;

- **Visual:**
  - hallucinations, illusions; occipital, or temporal origin; complex hallucinations and a ‘tunnel vision’ phenomenon are exclusive to seizures of anteromedial temporal and occipitotemporal origin, whereas elementary hallucinations, illusions, and visual loss are common to both occipital and temporal lobe seizures;

- **Auditory:**
  - may indicate an origin in the superior temporal gyrus;

- **Olfactory:**
  - parosmia may occur in seizures of medial temporal lobe origin (uncus; uncinate fits);

- **Gustatory;**
- **Autonomic;**
- **Abdominal:**
  - rising epigastric sensation (visceral aura) of temporal lobe epilepsy;

- **Psychic:**
  - complex hallucinations or illusions that usually affect different senses, e.g. distortions of familiarity such as *deja vu* or *jamais vu* auras of
Automatic Obedience

Automatic obedience may be seen in startle syndromes such as the jumping Frenchmen of Maine, latah, and myriachit, when a sudden shout of, for example, 'jump' is followed by a jump. These are sometimes known as the startle-automatic obedience syndromes. Although initially classified (by Gilles de la Tourette) with tic syndromes, there are clear clinical and pathophysiological differences.

Reference

Cross Reference
Tic

Automatic Writing Behaviour

Automatic writing behaviour is a form of increased writing activity. It has been suggested that it should refer specifically to a permanently present or elicitable, compulsive, iterative and not necessarily complete, written reproduction of visually or orally perceived messages (cf. hypergraphia). This is characterized as a particular, sometimes isolated, form of utilization behaviour in which the inhibitory functions of the frontal lobes are suppressed.

Reference

Cross References
Hypergraphia; Utilization behaviour

Automatism

Automatisms are complex motor movements occurring in complex motor seizures, which resemble natural movements but occur in an inappropriate setting. These may occur during a state of impaired consciousness during or shortly after an epileptic seizure. There is usually amnesia for the event.

Automatisms occur in about one-third of patients with complex partial seizures, most commonly those of temporal or frontal lobe origin. Although
there are qualitative differences between the automatisms seen in seizures arising from these sites, they are not of sufficient specificity to be of reliable diagnostic value; bizarre automatisms are more likely to be frontal.

Automatisms may take various forms:

- **Oro-facial movements:**
  - for example, lip smacking, chewing and swallowing movements, salivation (especially temporal lobe origin).

- **Gestural:**
  - hand fumbling, foot shuffling, tidying, or more complex actions such as undressing; upper limb movements are said to be more suggestive of temporal lobe origin, lower limb movements (kicking, cycling) of frontal lobe origin; pelvic thrusting (may also be seen in pseudo-seizures).

- **Ambulatory:**
  - walking or running around (cursive seizures); prolonged wandering may be termed fugue or poriomania.

- **Emotional:**
  - laughing and, more rarely, crying (gelastic and dacrystic seizures, respectively, although crying may also be a feature of non-epileptic seizures), fear, anger.

- **Verbal:**
  - humming, whistling, grunting, speaking incoherently; vocalization is common in frontal lobe automatisms.

Automatic behaviour and fugue-like states may also occur in the context of narcolepsy and must be differentiated from the automatisms of complex partial seizures on the basis of history, examination, and EEG.

**References**


**Cross References**

Absence; Aura; Pelvic thrusting; Poriomania; Seizure

**Autophony**

The perception of the reverberation of one's own voice, which occurs with external or middle, but not inner, ear disease.

**Autoscopy**

Autoscopy (literally ‘seeing oneself’) is a visual hallucination of one’s own face, sometimes with upper body or entire body, likened to seeing oneself in a mirror (hence mirror hallucination). The hallucinated image is a mirror image, i.e. shows left–right reversal as in a mirror image. Unlike heautoscopy, there is a coincidence of egocentric and body-centred perspectives. Autoscopy may be associated with parieto-occipital space-occupying lesions, epilepsy, and migraine.
Autotopagnosia

References

Cross References
Hallucination; Heautoscopy

**Autotopagnosia**

Autotopagnosia, or somatotopagnosia, is a rare disorder of body schema characterized by inability to identify parts of the body, either to verbal command or by imitation; this is sometimes localized but at worst involves all parts of the body. This may be a form of category-specific anomia with maximum difficulty for naming body parts or one feature of anosognosia. Finger agnosia and right–left disorientation are partial forms of autotopagnosia, all of which are most often seen following cerebrovascular events involving the left parietal area.

Reference

Cross References
Agnosia; Anosognosia; Finger agnosia; Gerstmann syndrome; Right–left disorientation; Somatoparaphrenia
Babinski’s Sign (1)

Babinski’s sign is a polysynaptic cutaneous reflex consisting of an extensor movement (dorsiflexion) of the big toe on eliciting the plantar response, due to contraction of extensor hallucis longus. There may be in addition fanning (abduction) of the other toes (fan sign; signe de l’éventail) but this is neither necessary nor sufficient for Babinski’s sign to be judged present. There may be simultaneous contraction of other limb flexor muscles, consistent with the notion that Babinski’s sign forms part of a flexion synergy (withdrawal) of the leg. The use of the term ‘negative Babinski sign’ to indicate the normal finding of a down-going (flexor; plantar flexion) big toe is incorrect, ‘flexor plantar response’ being the appropriate description. The plantar response is most commonly performed by stroking the sole of the foot, although many other variants are described (e.g. Chaddock’s sign, Gordon’s sign, Oppenheim’s sign).

Babinski’s sign is a normal finding in infants with immature (unmyelinated) corticospinal tracts; persistence beyond 3 years of age, or re-emergence in adult life, is pathological. In this context, Babinski’s sign is considered a reliable (‘hard’) sign of corticospinal (pyramidal) tract dysfunction (upper motor neurone pathology) and may coexist with other signs of upper motor neurone dysfunction (e.g. weakness in a so-called pyramidal distribution, spasticity, hyperreflexia). However, if weakness of extensor hallucis longus is one of the features of upper motor neurone dysfunction, or from any other cause, Babinski’s sign may be unexpectedly absent although anticipated on clinical grounds. Other causes of Babinski’s sign include hepatic coma, postepileptic seizure, deep sleep following prolonged induced wakefulness, and cataplectic attack, hence it is not necessarily a consequence of a permanent and irreversible lesion of the pyramidal tracts.

In the presence of extrapyramidal signs, it is important to distinguish Babinski’s sign, a ‘pyramidal sign’, from a striatal toe (spontaneous up going plantar).

References


Cross References

Chaddock’s sign; Gordon’s sign; Hyperreflexia; Oppenheim’s sign; Parkinsonism; Plantar response; Spasticity; Striatal toe; Upper motor neurone (UMN) syndrome; Weakness
Babinski’s Sign (2)
Babinski (1905) described the paradoxical elevation of the eyebrow in hemifacial spasm as orbicularis oris contracts and the eye closes, a synkinesis which is not reproducible by will. This observation indicated to Babinski the peripheral (facial nerve) origin of hemifacial spasm. It may assist in differentiating hemifacial spasm from other craniofacial movement disorders.

Reference

Cross Reference
Hemifacial spasm

Babinski’s Trunk–Thigh Test
Babinski’s trunk–thigh test, also known as the ‘rising sign’, is suggested to be of use in distinguishing organic from functional paraplegia and hemiplegia (the abductor sign may also be of use in the former case, Hoover’s sign in the latter). The recumbent patient is asked to sit up with the arms folded on the front of the chest. In organic hemiplegia there is involuntary flexion of the paretic leg, which may automatically rise higher than the normal leg; in paraplegia both legs are involuntarily raised. In functional paraplegic weakness neither leg is raised, and in functional hemiplegia only the normal leg is raised.

Reference

Cross References
Abductor sign; Functional weakness and sensory disturbance; Hemiplegia; Hoover’s sign; Paraplegia

Bag of Worms
- see MYOKYMIA

Balaclava Helmet
A pattern of facial sensory loss resembling in distribution a balaclava helmet, involving the outer parts of the face but sparing the nose and mouth, may be seen with central brainstem lesions such as syringobulbia which progress upwards from the neck, such that the lowermost part of the spinal nucleus of the trigeminal nerve which serves the outer part of the face is involved whilst the upper part of the nucleus which serves the central part of the face is spared. This pattern of facial sensory impairment may also be known as onion peel or onion skin.

Cross Reference
Onion peel, Onion skin

Balint’s Syndrome
Balint’s syndrome, first described by a Hungarian neurologist in 1909, consists of:
- Simultanagnosia (q.v.; dorsal type):
  A constriction of visual attention, such that the patient is aware of only one object at a time; visual acuity is preserved, and patients can
recognize single objects placed directly in front of them; they are unable
to read or distinguish overlapping figures.

- **Spatial disorientation:**
  Loss of spatial reference and memory, leaving the patient ‘lost in space’.

- **Disorders of oculomotor function:**
  Specifically, visually guided eye movements (fixation, pursuit, saccades); Balint’s ‘psychic paralysis of gaze’, or ‘sticky fixation’, refers
to an inability to direct voluntary eye movements to visual targets,
despite a full range of eye movements; this has also been character-
ized as a form of oculomotor apraxia. Accurate eye movements may
be programmed by sound or touch. Loss of spontaneous blinking has
also been reported.

- **Optic ataxia:**
  A failure to grasp or touch an object under visual guidance.

Not all elements may be present; there may also be coexisting visual field
defects, hemispatial neglect, visual agnosia, or prosopagnosia.

Balint’s syndrome results from bilateral lesions of the parieto-occipital junc-
tion causing a functional disconnection between higher-order visual cortical
regions and the frontal eye fields, with sparing of the primary visual cortex.
Brain imaging, either structural (CT, MRI) or functional (SPECT, PET), may
demonstrate this bilateral damage, which is usually of vascular origin, for ex-
ample, due to watershed or border zone ischaemia or top-of-the-basilar syndrome.
Balint syndrome has also been reported as a migrainous phenomenon, following
traumatic brain injury and in association with Alzheimer’s disease, brain tumour
(butterfly glioma), radiation necrosis, progressive multifocal leucoencephalopa-
thy, Marchiafava–Bignami disease with pathology affecting the corpus callosum,
and X-linked adrenoleucodystrophy.

**References**
Husein M, Stein J. Rezso Balint and his most celebrated case. *Archives of
Rafal R. Bálint’s syndrome: a disorder of visual cognition. In: D’Esposito M
(ed.). *Neurological foundations of cognitive neuroscience*. Cambridge, MA: MIT

**Cross References**
Apraxia; Blinking; Ocular apraxia; Optic ataxia; Simultanagnosia

**Ballism, Ballismus**
Ballism or ballismus is a hyperkinetic involuntary movement disorder char-
acterized by wild, flinging, throwing movements of a limb. These movements
most usually involve one-half of the body (hemiballismus), although they may
sometimes involve a single extremity (monoballismus) or both halves of the
body (paraballismus). The movements are often continuous during wakefulness
but cease during sleep. Hemiballismus may be associated with limb hypotonia.
Clinical and pathophysiological studies suggest that ballism is a severe form
of chorea. It is most commonly associated with lesions of the contralateral
subthalamic nucleus.
Cross References
Chorea, Choreoathetosis; Hemiballismus; Hypotonia, Hypotonus

Bathing Suit Sensory Loss
- see SUSPENDED SENSORY LOSS

Battle’s Sign
Battle’s sign is a haematoma overlying the mastoid process, which indicates an underlying basilar skull fracture extending into the mastoid portion of the temporal bone. It appears 48–72 h after the trauma which causes the fracture.

Beevor’s Sign
Beevor’s sign is an upward movement of the umbilicus in a supine patient attempting either to flex the head onto the chest against resistance (e.g. the examiner’s hand) or performing a sit-up. It indicates a lesion causing rectus abdominis muscle weakness below the umbilicus. This may occur with a spinal lesion (e.g. tumour, syringomyelia) between T10 and T12 causing isolated weakness of the lower part of the muscle, or myopathies affecting abdominal muscles, particularly facioscapulohumeral muscular dystrophy. Lower cutaneous abdominal reflexes are also absent, having the same localizing value.

Downward movement of the umbilicus (‘inverted Beevor’s sign’) due to weakness of the upper part of rectus abdominis is less often seen.

References

Cross Reference
Abdominal reflexes

Belle Indifférence
La belle indifférence refers to a patient’s seeming lack of concern in the presence of serious symptoms. This was first defined in the context of ‘hysteria’, along with exaggerated emotional reactions, what might now be termed functional or somatoform illness. However, the sign is a poor discriminator against ‘organic’ illness. Some patients’ coping style is to make light of serious symptoms; they might be labelled stoical.

Patients with neuropathological lesions may also demonstrate a lack of concern for their disabilities, either due to a disorder of body schema (anosodiaphoria) or due to incongruence of mood (typically in frontal lobe syndromes, sometimes seen in multiple sclerosis).

Reference
Bell's Palsy
Bell's palsy is an idiopathic peripheral (lower motor neurone) facial weakness (prosopoplegia). It is thought to result from viral inflammation of the facial (VII) nerve. Other causes of lower motor neurone facial paresis may need to be excluded before a diagnosis of Bell's palsy can be made.

In the majority of patients with Bell's palsy (idiopathic facial paresis), spontaneous recovery occurs over 3 weeks to 2 months. Poorer prognosis is associated with older age (over 40 years) and if no recovery is seen within 4 weeks of onset. Meta-analyses suggest that steroids are associated with better outcome than no treatment, but that acyclovir alone has no benefit.

References

Bell's Phenomenon, Bell's Sign
Bell's phenomenon or sign is reflex upward, and slightly outward, deviation of the eyes in response to forced closure, or attempted closure, of the eyelids. This is a synkinesis of central origin involving superior rectus and inferior oblique muscles. It may be very evident in a patient with Bell's palsy (idiopathic facial nerve paralysis) attempting to close the paretic eyelid. The reflex indicates intact nuclear and infranuclear mechanisms of upward gaze, and hence that any defect of upgaze is supranuclear. However, in making this interpretation it should be remembered that perhaps 10–15% of the normal population do not show a Bell’s phenomenon.

Bell's phenomenon is usually absent in progressive supranuclear palsy and is only sometimes spared in Parinaud's syndrome.

Reference

Cross References
Bell's palsy; Gaze palsy; Parinaud's syndrome; Supranuclear gaze palsy; Synkinesia, Synkinesis

Benediction Hand
Median nerve lesions in the axilla or upper arm cause weakness in all median nerve innervated muscles, including flexor digitorum profundus. Thus
on attempting to make a fist, impaired flexion of the index and middle fingers, complete and partial, respectively, but with normal ring and little finger flexion (ulnar nerve mediated) results in a hand posture likened to that of a priest saying benediction (also sometimes known as Benedictine hand or orator’s hand).

**Cross References**
Claw hand; Simian hand

**Bent Spine Syndrome**
- see CAMPTOCORMIA

**Bielschowsky’s Sign, Bielschowsky’s Test**
Bielschowsky’s sign is head tilt towards the shoulder, typically towards the side contralateral to a trochlear (IV) nerve palsy. The intorsion of the unaffected eye brought about by the head tilt compensates for the double vision caused by the unopposed extorsion of the affected eye. Very occasionally, head tilt is paradoxical, i.e. towards the involved side: presumably the greater separation of images thus produced allows one of them to be ignored.

Bielschowsky’s (head tilt) test consists of the examiner tipping the patient’s head from shoulder to shoulder to see if this improves or exacerbates double vision, as will be the case when the head is, respectively, tilted away from or towards the affected side in a unilateral trochlear (IV) nerve lesion. The test is usually negative in a skew deviation causing vertical divergence of the eyes. This test may also be used as part of the assessment of vertical diplopia to see whether hypertropia changes with head tilt to left or right; increased hypertropia on left head tilt suggests a weak intortor of the left eye (superior rectus); increased hypertropia on right head tilt suggests a weak intortor of the right eye (superior oblique).

**Cross References**
Diplopia; Hypertropia; Skew deviation

**Binasal Hemianopia**
Of the hemianopic defects, binasal hemianopia, suggesting lateral compression of the chiasm, is less common than bitemporal hemianopia. Various causes are recorded including syphilis, glaucoma, drusen, and chronically raised intracranial pressure.

**Reference**

**Cross Reference**
Hemianopia

**Bitemporal Hemianopia**
Bitemporal hemianopia due to chiasmal compression, for example, by a pituitary lesion or craniopharyngioma, is probably the most common cause of a heteronymous hemianopia. Conditions mimicking bitemporal hemianopia include congenitally tilted discs, nasal sector retinitis pigmentosa, and papilloedema with greatly enlarged blind spots.
Blindsight

Cross References
Hemianopia; Visual field defects

Blepharoptosis
- see PTOSIS

Blepharospasm
Blepharospasm is a focal dystonia of the orbicularis oculi resulting in repeated involuntary forced eyelid closure, with failure of voluntary eye opening. Usually bilateral in origin, it may be sufficiently severe to result in functional blindness. The condition typically begins in the sixth decade of life and is more common in women than in men. Blepharospasm may occur in isolation (‘benign essential blepharospasm’), or in combination with other involuntary movements which may be dystonic (orobuccolingual dystonia or Meige syndrome; limb dystonia) or dyspraxic (eyelid apraxia), or in association with another neurological disorder such as Parkinson’s disease. Other examples of ‘secondary blepharospasm’ include drug therapy (neuroleptics, levodopa) and lesions of the brainstem and more rarely cerebellum and striatum. Like other forms of dystonia, blepharospasm may be relieved by sensory tricks (*geste antagoniste*), such as talking, yawning, singing, humming, or touching the eyelid. This feature is helpful in diagnosis. Blepharospasm may be aggravated by reading, watching television, and exposure to wind or bright light.

Blepharospasm is usually idiopathic but may be associated with lesions (usually infarction) of the rostral brainstem, diencephalon, and striatum; it has been occasionally reported with thalamic lesions. The pathophysiological mechanisms underlying blepharospasm are not understood, but may reflect dopaminergic pathway disruption causing disinhibition of brainstem reflexes. Local injections of botulinum toxin into orbicularis oculi are the treatment of choice, the majority of patients deriving benefit and requesting further injection. Failure to respond to botulinum toxin may be due to concurrent eyelid apraxia or dopaminergic therapy with levodopa.

References

Cross References
Blinking; Dystonia; Eyelid apraxia; Gaping; *Geste antagoniste*; Yawning

Blindsight
Blindsight describes a rare phenomenon in which patients with bilateral occipital lobe damage affecting the primary visual cortex are nonetheless able to discriminate certain visual events within their ‘blind’ fields, but are not aware of their ability to do so.

Reference
Cross Reference
Scotoma

Blind Spot
The blind spot is defined anatomically as the point on the retina at which axons from the retinal ganglion cells enter the optic nerve; since this area is devoid of photoreceptors there is a physiological blind spot. This area may be mapped clinically by confrontation with the examiner’s blind spot or mechanically. Minor enlargement of the blind spot is difficult to identify clinically, formal perimetry is needed in this situation.

Enlargement of the blind spot (peripapillary scotoma) is observed with raised intracranial pressure causing papilloedema: this may be helpful in differentiating papilloedema from other causes of disc swelling such as optic neuritis, in which a central scotoma is the most common field defect. Enlargement of the blind spot may also be a feature of peripapillary retinal disorders including big blind spot syndrome.

Cross References
Disc swelling; Papilloedema; Scotoma

Blinking
Involuntary blinking rate is decreased in idiopathic Parkinson’s disease (and may be improved by dopaminergic therapy) and in progressive supranuclear palsy (Steele–Richardson–Olszewski syndrome) where the rate may be <5/min. In contrast, blink rate is normal in multiple system atrophy and dopa-responsive dystonia, and increased in schizophrenia and postencephalitic parkinsonism. These disparate observations are not easily reconciled with the suggestion that blinking might be a marker of central dopaminergic activity.

Loss of spontaneous blinking has been reported in Balint’s syndrome. In patients with impaired consciousness, the presence of involuntary blinking implies an intact pontine reticular formation; absence suggests structural or metabolic dysfunction of the reticular formation. Blinking decreases in coma. Functional disorders may be accompanied by an increase in blinking.

Reference

Cross References
Balint’s syndrome; Blink reflex; Coma; Corneal reflex; Parkinsonism; Sighing; Yawning

Blink Reflex
The blink reflex consists of bilateral reflex contraction of the orbicularis oculi muscles. This may be induced by:

- Mechanical stimulus:
  Examples include percussion over the supraorbital ridge (glabellar tap reflex, Myerson’s sign, nasopalpebral reflex): this quickly habituates with repetitive stimulation in normal individuals; touching the cornea (corneal reflex); stroking the eyelashes in unconscious patients with closed eyes (‘eyelash reflex’).
• **Visual stimulus:**
  Sudden visual stimulus approaching the eyes (menace reflex, threat reflex, visuopalpebral reflex): the stimulus should be unexpected since the reflex can be voluntarily suppressed; failure to respond to a stimulus moving into the temporal field of vision may indicate a hemianopic field defect in patients unable to comply with standard confrontation visual field testing. Care should be taken to avoid generating air currents with the hand movement as this may stimulate the corneal reflex which may simulate the visuopalpebral reflex. It is probable that this reflex requires cortical processing: it is lost in persistent vegetative states. Loss of this reflex may occur in Balint’s syndrome, ascribed to inability to recognize the nearness of the threatening object.

• **Acoustic stimulus:**
  Sudden loud sounds (acousticopalpebral reflex).

The final common (efferent) pathway for these responses is the facial nerve nucleus and facial (VII) nerve, the afferent limbs being the trigeminal (V), optic (II), and auditory (VIII) nerves, respectively.

Electrophysiological study of the blink reflex may demonstrate peripheral or central lesions of the trigeminal (V) nerve or facial (VII) nerve (afferent and efferent pathways, respectively). It has been reported that in the evaluation of sensory neuronopathy the finding of an abnormal blink reflex favours a non-paraneoplastic aetiology, since the blink reflex is normal in paraneoplastic sensory neuronopathies.

**References**

**Cross References**
Balint’s syndrome; Blinking; Corneal reflex; Glabellar tap reflex

**Body Part as Object**
In this phenomenon, apraxic patients use a body part when asked to pantomime certain actions, such as using the palm when asked to demonstrate the use of a hair brush or comb, or fingers when asked to demonstrate use of scissors or a toothbrush.

**References**

**Cross References**
Apraxia; Parapraxia, Parapraxis
**Bon-Bon Sign**
Involuntary pushing of the tongue against the inside of the cheek, the ‘bon-bon sign’, is said to be typical of the stereotypic oro-lingual movements of tardive dyskinesia, along with chewing and smacking of the mouth and lips, and rolling of the tongue in the mouth. These signs may help to distinguish tardive dyskinesia from chorea, although periodic protrusion of the tongue (flycatcher, trombone tongue) is common to both.

**Cross References**
Buccolingual syndrome; Chorea, Choreoathetosis; Trombone tongue

**Bouche de Tapir**
Patients with facioscapulohumeral (FSH) muscular dystrophy have a peculiar and characteristic facies, with puckering of the lips when attempting to whistle. The pouting quality of the mouth, unlike that seen with other types of bilateral (neurogenic) facial weakness, has been likened to the face of the tapir (*Tapirus* sp.).

**Cross Reference**
Facial paresis

**Bovine Cough**
A bovine cough lacks the explosive character of a normal voluntary cough. It may result from injury to the distal part of the vagus nerve, particularly the recurrent laryngeal branches which innervate all the muscles of the larynx (with the exception of cricothyroid) with resultant vocal cord paresis. Because of its longer intrathoracic course, the left recurrent laryngeal nerve is more often involved. A bovine cough may be heard in patients with tumours of the upper lobes of the lung (Pancoast tumour) due to recurrent laryngeal nerve palsy. Bovine cough may also result from any cause of bulbar weakness, such as motor neurone disease, Guillain–Barré syndrome, and bulbar myopathies.

**Reference**

**Cross References**
Bulbar palsy; Diplophonia; *Signe de rideau*

**Bradykinesia**
Bradykinesia is a slowness in the initiation and performance of voluntary movements in the absence of weakness and is one of the typical signs of parkinsonian syndromes, in which situation it is often accompanied by difficulty in the initiation of movement (akinesia, hypokinesia) and reduced amplitude of movement (hypometria) which may increase with rapid repetitive movements (fatigue). It may be overcome by reflexive movements or in moments of intense emotion (*kinesis paradoxica*).

Bradykinesia in parkinsonian syndromes reflects dopamine depletion in the basal ganglia. It may be improved by levodopa and dopaminergic agonists, less so by anticholinergic agents.
Slowness of voluntary movement may also be seen with psychomotor retardation, frontal lobe lesions producing abulia, and in the condition of obsessive slowness.

**Reference**

**Cross References**
Abulia; Akinesia; Fatigue; Hypokinesia; Hypometria; *Kinesis paradoxica*; Parkinsonism; Psychomotor retardation

**Bradylalia**
Bradylalia is slowness of speech, typically seen in the frontal–subcortical types of cognitive impairment, with or without extrapyramidal features, or in depression.

**Cross References**
Palilalia; Tachylalia

**Bradyphrenia**
Bradyphrenia is a slowness of thought, typically seen in the frontal–subcortical types of cognitive impairment, e.g. progressive supranuclear palsy, vascular dementia, and Huntington’s disease. Such patients typically answer questions correctly but with long response times.

**Cross References**
Abulia; Dementia

**Bragard’s Test**
- see LASÈGUE’S SIGN

**Broca’s Aphasia**
Broca’s aphasia is the classic ‘expressive aphasia’, in distinction to the ‘receptive aphasia’ of Wernicke; however, there are problems with this simple classification, since Broca’s aphasics may show comprehension problems with complex material, particularly in relation to syntax. Considering each of the features suggested for the clinical classification of aphasias (see Aphasia), Broca’s aphasia is characterized by:

- **Fluency**: slow, laboured, effortful speech (non-fluent) with phonemic paraphasias, agrammatism, and aprosody; the patient knows what s/he wants to say and usually recognizes the paraphasic errors (i.e. patients can ‘self-monitor’);
- **Comprehension**: comprehension for simple material is preserved, but there may be problems with more complex syntax;
- **Repetition**: impaired;
- **Naming**: impaired (anomia, dysnomia); may be aided by phonemic or contextual cueing (cf. Wernicke’s aphasia);
- **Reading**: alexia with laboured oral reading, especially of function words and verb inflections. Silent reading may also be impaired (deep dyslexia) as reflected by poor text comprehension;
- **Writing**: similarly affected.
Aphemia was the name originally given by Broca to the language disorder subsequently named ‘Broca’s aphasia’. The term alalia was also once used. The terms ‘small Broca’s aphasia’, ‘mini-Broca’s aphasia’, and ‘Broca’s area aphasia’ have been reserved for a more circumscribed clinical and neuroanatomical deficit than Broca’s aphasia, wherein the damage is restricted to Broca’s area or its subjacent white matter. There is a mild and transient aphasia or anomia which may share some of the characteristics of aphemia phonetic disintegration (i.e. a motor disorder of speech production with preserved comprehension of spoken and written language).

The syndrome of Broca’s aphasia may emerge during recovery from a global aphasia. Broca’s aphasia is sometimes associated with a right hemiparesis, especially affecting the arm and face; there may also be bucco-lingual-facial dyspraxia. Depression may be a concurrent feature.

Classically Broca’s aphasia is associated with a vascular lesion of the third frontal gyrus in the inferior frontal lobe (Broca’s area), but in practice such a circumscribed lesion is seldom seen. More commonly there is infarction in the perisylvian region affecting the insula and operculum (Brodmann areas 44 and 45), which may include underlying white matter and the basal ganglia (territory of the superior branch of the middle cerebral artery).

References

Cross References
Agrammatism; Agraphia; Alalia; Alexia; Aphasia; Aphemia; Aprosodia, Aprosody; Paraphasia; Recurrent utterances; Wernicke’s aphasia

Brown-Séquard Syndrome
The Brown-Séquard syndrome is the consequence of anatomical or, more usually, functional hemisection of the spinal cord (spinal hemisection syndrome), producing the following pattern of clinical findings:

- **Motor:**
  Ipsilateral spastic weakness, due to corticospinal tract involvement;
  Segmental lower motor neurone signs at the level of the lesion, due to root and/or anterior horn cell involvement.

- **Sensory:**
  A dissociated sensory loss, i.e.:
  Ipsilateral loss of proprioception, due to dorsal column involvement;
  Contralateral loss of pain and temperature sensation, due to crossed spinothalamic tract involvement.

Spinal cord lesions producing this syndrome may be either extramedullary (e.g. prolapsed cervical intervertebral disc, extrinsic spinal cord tumour) or intramedullary (e.g. multiple sclerosis, intrinsic spinal cord tumour, myelitis, radiation-induced myelopathy); the former group is said to be the more common cause.
References

Cross References
Dissociated sensory loss; Myelopathy; Proprioception; Spasticity; Weakness

Brudzinski’s (Neck) Sign
Brudzinski described a number of signs, but the one most often used in clinical practice is the neck sign, which is sometimes evident in cases of meningeal irritation, for example, due to meningitis. Passive flexion of the neck to bring the head onto the chest is accompanied by flexion of the thighs and legs. As with nuchal rigidity and Kernig’s sign, Brudzinski’s sign may be absent in elderly or immunosuppressed patients with meningeal irritation.

Reference

Cross References
Kernig’s sign; Meningism; Nuchal rigidity

Brueghel’s Syndrome
Brueghel’s syndrome [NB some texts give ‘Breughel’s’ syndrome] is the name given to a dystonia of the motor trigeminal nerve causing gaping or involuntary opening of the mouth, so named after Brueghel’s painting De Gaper of 1558, thought to illustrate a typical case. Additional features may include paroxysmal hyperpnoea and upbeating nystagmus. Brueghel’s syndrome should be distinguished from other syndromes of cranial dystonia featuring blepharospasm and oromandibular dystonia, better termed Meige’s syndrome.

Reference

Cross References
Blepharospasm; Dystonia

Bruit
Bruit arise from turbulent blood flow causing arterial wall vibrations which are audible at the body surface with the unassisted ear or with a stethoscope (diaphragm rather than bell, better for detecting higher frequency sounds). They are associated with stenotic vessels or with fistulae where there is arteriovenous shunting of blood. Dependent on the clinical indication, various sites may be auscultated: eye for orbital bruit in carotico-cavernous fistula; head for bruit of AV fistula; but probably the most frequently auscultated region is the carotid bifurcation, high up under the angle of the jaw, in individuals thought to have had a transient ischaemic attack or ischaemic stroke. Examination for carotid bruits in asymptomatic individuals is probably best avoided, other than in the clinical trial
setting, since the optimal management of asymptomatic carotid artery stenosis has yet to be fully defined.

**Reference**

**Brushfield Spots**
Brushfield spots are small grey-white specks of depigmentation that can be seen in the irides of some (90%) patients with Down’s syndrome; they may also occur in normal individuals.

**Bruxism**
Bruxism is forcible grinding or gnashing of the teeth. This is common in children and is said to occur in 5–20% of the population during non-REM sleep (a parasomnia). Masseter hypertrophy may become apparent in persistent grinders. Bruxism may also occur in encephalopathic disorders (e.g. hepatic encephalopathy) and occasionally in disorders of the basal ganglia (multiple system atrophy, basal ganglia infarcts). Dysfunction of efferent and/or afferent thalamic and striatopallidal tracts has been suggested as the neural substrate. If necessary, a rubber gum shield or bite may be worn in the mouth to protect the teeth. Botulinum toxin injections have also been tried.

**Reference**

**Cross References**
Encephalopathy; Masseter hypertrophy

**Buccofacial Dyspraxia**
- see ORO-FACIAL DYSPRAXIA

**Buccolingual Syndrome**
This is a form of tardive dyskinesia that involves involuntary movements of the facial muscles and protrusion of the tongue.

**Cross References**
‘Bon-bon sign’; Dyskinesia

**Bulbar Palsy**
Bulbar palsy is weakness of bulbar musculature of lower motor neurone origin. This may be differentiated clinically from bulbar weakness of upper motor neurone origin (pseudobulbar palsy).

Clinical features of bulbar palsy include
- Dysarthria of flaccid/nasal type;
- Dysphonia;
- Dysphagia, often with nasal regurgitation;
- Weak (‘bovine’) cough; risk of aspiration;
- +/- wasted, fasciculating tongue;
- +/- absent jaw jerk;
- +/- absent gag reflex.
Bulbar palsy is usually neurogenic. Recognized causes include

- **Brainstem disorders affecting cranial nerve motor nuclei (intrinsic):**
  - Motor neurone disease (which may also cause a pseudobulbar palsy);
  - Poliomyelitis;
  - Glioma;
  - Syringobulbia.

- **Cranial nerve lesions outside the brainstem (there may be associated sensory signs):**
  - Infiltration by carcinoma, granuloma.

- **Neuromuscular junction transmission defect:**
  - Myasthenia gravis.

A myogenic bulbar palsy may be seen in oculopharyngeal muscular dystrophy, inclusion body myositis, and polymyositis.

**Cross References**
- Bovine cough; Dysarthria; Dysphagia; Dysphonia; Fasciculation; Gag reflex; Jaw jerk; Lower motor neurone (LMN) syndrome; Pseudobulbar palsy; Upper motor neurone (UMN) syndrome

**Bulbocavernous Reflex**
A test of the integrity of the S2, S3, and S4 spinal roots, looking for contraction of the anal sphincter (may be felt with a gloved finger in the rectum) when squeezing the glans penis or clitoris. The reflex may be abolished in lesions of the cauda equina.

**Cross References**
- Cauda equina syndrome; Reflexes

**Buphthalmos**
Buphthalmos, literally ox-eye, consists of a large and bulging eye caused by raised intraocular pressure due to congenital or secondary glaucoma. This is one of the ophthalmological features of Sturge–Weber syndrome.

**Butt-First Manoeuvre**
- see Gowers’ SIGN
Cacogeusia
Sensation of a disagreeable taste, often associated with parosmia.

Cacosmia
- see PAROSMIA

Calf Head Sign
A consistent pattern of muscle enlargement or wasting, described as ‘calf heads on a trophy’, has been observed in Miyoshi-type dysferlinopathy when the arms are raised with shoulder abducted and elbows flexed to 90°. Diamond on quadriceps sign may also be seen in dysferlinopathies.

Reference

Cross Reference
Diamond on quadriceps sign

Calf Hypertrophy
Calf enlargement has many causes; it may reflect true hypertrophy (enlargement of muscle fibres) or, more commonly, pseudohypertrophy, due to infiltration with tissue elements other than muscle.

Hypertrophy may be due to neuromuscular disorders producing

- chronic partial denervation, e.g.:
  - radiculopathy;
  - peripheral neuropathy;
  - spinal muscular atrophy;
  - following paralytic poliomyelitis.

- continuous muscle activity, e.g.:
  - myotonia congenital;
  - Isaac’s syndrome (neuromyotonia);
  - generalized myokymia.

Calf (and other muscle) hypertrophy is also a feature of limb girdle muscular dystrophy type 2I.

Calf pseudohypertrophy may be due to:

- Dystrophinopathies (Duchenne muscular dystrophy, Becker dystrophy), due to excess connective tissue;
- Infection/inflammation: myositis;
- Infiltration: amyloidosis, tumour, cysticercosis.
Caloric Testing

Caloric tests examine the vestibulo-ocular reflexes (VOR). They are mainly used in two circumstances: to identify vestibular pathology in the assessment of dizziness/vertigo when clinical tests of VOR are unhelpful and to assess brainstem integrity in coma.

Each labyrinth may be separately assessed by irrigating each outer ear. Head flexion to 30° above the horizontal allows maximum stimulation of the horizontal semicircular canals, whereas 60° below horizontal maximally stimulates the lateral semicircular canals. Water 7°C above and below body temperature (i.e. 30°C and 44°C) is used, applied for 30–40 s. Induced nystagmus is then timed both with and without visual fixation (in the dark, Frenzel glasses). This method is cheap but has poor patient acceptability.

Normally, the eyes show conjugate deviation towards the ear irrigated with cold water, with corrective nystagmus in the opposite direction; with warm water the opposite pattern is seen. (The direction of nystagmus may thus be recalled by the mnemonic COWS: cold opposite, warm same.) Dysconjugate responses suggest brainstem damage or depression. A reduced duration of induced nystagmus is seen with canal paresis; enhancement of the nystagmus with removal of visual fixation suggests this is peripheral in origin (labyrinthine, vestibulocochlear nerve), whereas no enhancement suggests a central lesion.

In coma the deviation may be present but without corrective saccades, even at a time when the oculocephalic responses elicited by the doll’s head manoeuvre are lost. As coma deepens even the caloric reflexes are lost as brainstem involvement progresses.

Reference

Cross References
Coma; Nystagmus; Oculocephalic response; Vertigo; Vestibulo-ocular reflexes

Camptocormia

Camptocormia, or ‘bent spine syndrome’, was first described as a psychiatric phenomenon in men facing armed conflict (a ‘war neurosis’). It has subsequently been realized that reducible lumbar kyphosis may also result from neurological disorders, including muscle disease (paravertebral myopathy, nemaline myopathy), Parkinson’s disease, dystonia, motor neurone disease, and, possibly, as a paraneoplastic phenomenon. Cases with associated lenticular (putaminal) lesions have also been described.
Camptocormia may be related in some instances to dropped head syndrome.

References

Cross References
Dropped head syndrome; Dystonia

Camptodactyly
Camptodactyly, literally ‘bent finger’, is a flexion deformity at the proximal interphalangeal joint, especially affecting the little fingers; this may be unilateral or bilateral. A distinction is sometimes drawn between camptodactyly and streblodactyly: in the latter, several fingers are affected by flexion contractures (strebl = twisted, crooked), but it is not clear whether the two conditions overlap or are separate. The term streblomicrodactyly has sometimes been used to designate isolated crooked little fingers. Camptodactyly is not accompanied by any sensory or motor signs. The condition may be familial and is more common in women. Camptodactyly may occur as part of a developmental disorder with other dysmorphic features or in isolation.

It is important to differentiate camptodactyly, a non-neurogenic cause of clawing, from neurological diagnoses such as:

- Ulnar neuropathy;
- C8/T1 radiculopathy;
- Cervical rib;
- Syringomyelia.

Awareness of the condition is important to avoid unnecessary neurological investigation.

Reference

Cross Reference
Claw hand

Capgras Syndrome
This is one of the classical delusional syndromes of psychiatry, in which patients recognize a close family relative, or other loved object, but believe them to be have been replaced by an exact alien or ‘double’ (illusion of doubles). Initially described in patients with psychiatric disorders, it may also occur in traumatic, metabolic, and neurodegenerative disorders (e.g. Alzheimer’s disease, dementia with Lewy bodies). Neurologists have encompassed this phenomenon under the term reduplicative paramnesia. Some believe this syndrome to be the ‘mirror image’ of prosopagnosia, in which faces are not recognized but emotional significance is. Capgras syndrome may be envisaged as a Geschwindian disconnection syndrome, in which the visual recognition system is disconnected from the limbic system, hence faces can be recognized but no emotional significance ascribed to them.
References

Cross References
Cotard’s syndrome; Disconnection syndromes; Prosopagnosia; Reduplicative paramnesia

Carphologia
Carphologia, or floccillation, is an aimless plucking at clothing, as if picking off pieces of thread. This may sometimes be seen in psychiatric illness, delirium, Alzheimer’s disease, or vascular dementia particularly affecting the frontal lobe. Some have characterized carphologia as a form of akathisia.

Reference

Cross References
Akathisia; Delirium; Dementia

Carpopedal Spasm
- see *MAIN D’ACCOUCHEUR*

Catalepsy
This term has been used to describe increased muscle tone, leading to the assumption of fixed postures which may be held for long periods of time without apparent fatigue; it may be possible for the examiner to position an extremity into any posture, in which it then remains for some time. Clearly, this term is cognate with or overlaps with waxy flexibility which is a feature of catatonic syndromes. Catalepsy may be feigned (see Dr Arthur Conan Doyle’s story of The Resident Patient in *The Memoirs of Sherlock Holmes*, first published in 1894).

Catalepsy should not be confused with the term cataplexy, a syndrome in which muscle tone is transiently lost.

Reference

Cross Reference
Cataplexy; Catatonia

Cataplexy
Cataplexy is a sudden loss of limb tone which may lead to falls (drop attacks) without loss of consciousness, usually lasting less than 1 min. Attacks may be precipitated by strong emotion (laughter, anger, embarrassment, surprise). Sagging of the jaw and face may occur, as may twitching around the face or eyelids. During an attack there is electrical silence in antigravity muscles, which are consequently hypotonic, and transient areflexia. Rarely status cataplecticus may develop, particularly after withdrawal of tricyclic antidepressant medication.
Cataplexy may occur as part of the narcoleptic syndrome of excessive and inappropriate daytime somnolence, hypnagogic hallucinations, and sleep paralysis (Gélineau’s original description of narcolepsy in 1877 included an account of ‘astasia’ which corresponds to cataplexy). Symptomatic cataplexy occurs in certain neurological diseases including brainstem lesions, Von Economo’s disease (postencephalitic parkinsonism), Niemann–Pick disease type C, and Norrie’s disease.

Therapeutic options for cataplexy include tricyclic antidepressants such as protriptyline, imipramine, and clomipramine; serotonin-reuptake inhibitors such as fluoxetine; and noradrenaline and serotonin-reuptake inhibitors such as venlafaxine.

References

Cross References
Areflexia; Hypersomnolence; Hypotonia, Hypotonus

Catathrenia
Catathrenia is expiratory groaning during sleep, especially its later stages. Although sufferers are unaware of the condition, it does alarm relatives and bed partners. There are no associated neurological abnormalities and no identified neurological or otorhinolaryngological cause. Catathrenia is categorized with the parasomnias in the International Classification of Sleep Disorders (ICSD2, 2005).

Reference

Catatonia
Catatonia is a clinical syndrome, first described by Kahlbaum (1874), characterized by a state of unresponsiveness but with maintained, immobile, body posture (sitting, standing; cf. stupor), mutism, and refusal to eat or drink, with or without staring, grimacing, limb rigidity, maintained abnormal postures (waxy flexibility or *flexibilitas cerea*), negativism, echophenomena (imitation behaviour), stereotypy, and urinary incontinence or retention. After recovery patients are often able to recall events which occurred during the catatonic state (cf. stupor). ‘Lethal catatonia’, in which accompanying fever and collapse lead to death, was described in the 1930s and seems to resemble neuroleptic malignant syndrome; the name ‘malignant catatonia’ has been proposed for this syndrome. Catatonia may be confused clinically with abulia.

Kraepelin classified catatonia as a subtype of schizophrenia but most catatonic patients in fact suffer a mood or affective disorder. Furthermore, although initially thought to be exclusively a feature of psychiatric disease, catatonia is now recognized as a feature of structural or metabolic brain disease (the original account contains descriptions suggestive of extrapyramidal disease):
- Psychiatric disorders:
  - Manic-depressive illness;
  - Schizophrenia.

- Neurological disorders:
  - Cerebrovascular disease (posterior circulation);
  - Tumours (especially around third ventricle, corpus callosum);
  - Head trauma;
  - Encephalitis;
  - Neurosyphilis;
  - Extrapyramidal disorders;
  - Epilepsy.

- Systemic illnesses:
  - Endocrine: hyperthyroidism, Addison’s disease, Cushing’s disease, diabetic ketoacidosis;
  - Metabolic: uraemia, hypercalcaemia, hepatic encephalopathy;
  - Others: systemic lupus erythematosus.

Various subtypes of catatonia are enumerated by some authorities, including

- Retarded catatonia (Kahlbaum’s syndrome);
- Excited catatonia (manic delirium, Bell’s mania);
- Malignant catatonia, lethal catatonia: also encompasses the neuroleptic malignant syndrome and the serotonin syndrome;
- Periodic catatonia.

Catatonia of psychiatric origin often responds to lorazepam; there are also advocates of ECT.

References

Cross References
Abulia; Akinetic mutism; Imitation behaviour; Mutism; Negativism; Rigidity; Stereotypy; Stupor

Cauda Equina Syndrome
A cauda equina syndrome results from pathological processes affecting the spinal roots below the termination of the spinal cord around L1/L2, hence it is a syndrome of multiple radiculopathies. Depending on precisely which roots are affected, this may produce symmetrical or asymmetrical sensory impairment in the buttocks (saddle anaesthesia; sacral anaesthesia) and the backs of the thighs, radicular pain, and lower motor neurone type weakness of the foot and/or toes (even a flail foot). Weakness of hip flexion (L1) does not occur, and
Central Scotoma, Centrocaecal Scotoma

this may be useful in differentiating a cauda equina syndrome from a conus lesion which may otherwise produce similar features. Sphincters may also be involved, resulting in incontinence, or, in the case of large central disc herniation at L4/L5 or L5/S1, acute urinary retention. Causes of a cauda equina syndrome include

- Central disc herniation;
- Tumour: primary (ependymoma, meningioma, Schwannoma), metastasis;
- Haematoma;
- Abscess;
- Lumbosacral fracture;
- Inflammatory disease, e.g. sarcoidosis (rare);
- Ankylosing spondylitis (rare).

The syndrome needs to be considered in any patient with acute (or acute-on-chronic) low back pain, radiation of pain to the legs, altered perineal sensation, and altered bladder function. Missed diagnosis of acute lumbar disc herniation may be costly, from the point of view of both clinical outcome and resultant litigation.

References

Cross References
Bulbocavernous reflex; Foot drop; Incontinence; Radiculopathy; Urinary retention

Central Scotoma, Centrocaecal Scotoma

These visual field defects are typical of retinal or optic nerve pathology. They may be mapped by confrontation testing or automatically.

- **Central scotoma:**
  Field defect occupying the macula, due to involvement of the macula or the papillomacular bundle; this is the typical (but not exclusive) finding in optic neuritis, but may also be seen with disease of the macula, optic nerve compression, and Leber’s hereditary optic neuropathy. Examination for a concurrent contralateral superior temporal defect should be undertaken: such junctional scotomas may be seen with lesions at the anterior angle of the chiasm.

- **Centrocaecal or caecocentral scotoma:**
  Field defect involving both the macula and the blind spot; seen in optic nerve disease, such as Leber’s hereditary optic neuropathy, toxic neuropathy, or nutritional optic neuropathy (said to be typical of vitamin B$_{12}$ deficiency optic neuropathy), sometimes in optic neuritis.
Cerebellar Syndromes

Differing clinical pictures may be seen with pathology in different parts of the cerebellum. Broadly speaking, a midline cerebellar syndrome (involving the vermis) may be distinguished from a hemispheric cerebellar syndrome (involving the hemispheres). Their clinical characteristics are as follows:

- **Midline cerebellar syndrome:** Gait ataxia but with little or no limb ataxia, hypotonia, or nystagmus (because the vestibulocerebellum is spared), or dysarthria; causes include alcoholic cerebellar degeneration, tumour of the midline (e.g. medulloblastoma), paraneoplastic cerebellar degeneration.

- **Hemispheric cerebellar syndrome:** Limb ataxia (e.g. ataxia on finger–nose and/or heel–shin testing), dysdiadochokinesia, dysmetria, dysarthria, nystagmus; usual causes are infarcts, haemorrhages, demyelination, and tumours.

- **Pancerebellar syndrome:** Affecting all parts of the cerebellum and showing a combination of the above signs (e.g. cerebellar degenerations).

Reference

Cross References
Asynergia; Ataxia; Dysarthria; Dysdiadochokinesia; Dysmetria; Hemiataxia; Hypotonia, Hypotonus; Nystagmus

Chaddock’s Sign
Chaddock’s sign, or the external malleolar sign, is a variant method for eliciting the plantar response, by application of a stimulus in a circular direction around the external malleolus, or the lateral aspect of the foot, moving from heel to little toe. Extension of the hallux (upgoing plantar response, Babinski’s sign) is pathological, indicating corticospinal tract (upper motor neurone) pathology. The development of Babinski’s sign always predates that of Chaddock’s sign.

References

Cross References
Babinski’s sign (1); Gordon’s sign; Oppenheim’s sign; Plantar response; Upper motor neurone (UMN) syndrome

Charcot Joint
Charcot joint, or neuropathic joint, describes a destructive arthropathy seen following repeated injury to an anaesthetic joint in patients with impaired or absent
pain sensation. There is trophic change, with progressive destruction of articular surfaces with disintegration and reorganization of joint structure. Although the destruction is painless, the Charcot joint itself may be painful. There may be concurrent skin ulceration.

Charcot joints were originally described in the context of tabes dorsalis (knees, shoulders, elbows, hips, ankles) but they may also be seen in:

- Syringomyelia (elbow);
- Hereditary sensory (and autonomic) neuropathies (HSAN, ‘congenital insensitivity to pain’; ankles);
- Leprosy;
- Diabetes mellitus.

Reference

Cross References
Analgesia; Main succulente

Charles Bonnet Syndrome
Described by the Swiss naturalist and philosopher Charles Bonnet in 1760, this syndrome consists of well-formed (complex), elaborated, and often stereotyped visual hallucinations, of variable frequency and duration, in a partially sighted (usually elderly) individual who has insight into their unreality. Hallucinations may disappear on eye closure. Predisposing visual disorders include cataract, macular degeneration, and glaucoma. There are no other features of psychosis or neurological disease such as dementia.

The pathogenesis of the visual hallucinations is uncertain. Reduced stimulation of the visual system leading to increased cortical hyperexcitability is one possible explanation (the deafferentation hypothesis), although the syndrome may occasionally occur in people with normal vision. Functional magnetic resonance imaging suggests ongoing cerebral activity in ventral extrastriate visual cortex.

Treatment consists primarily of reassurance. Pharmacological treatment with atypical antipsychotics or anticonvulsants may be tried but there is no secure evidence base.

References


**Cross References**

Hallucination; Pseudohallucination

**Chasm**
- see YAWNING

**Cheiro-Oral Syndrome**
- see PSEUDORADICULAR SYNDROME

**Cherry Red Spot at the Macula**
The appearance of a ‘cherry red spot at the macula’, caused by the contrast of a red macula against retinal pallor, occurs in a number of metabolic storage disorders, including

- Sialidosis (type I = cherry red spot–myoclonus syndrome);
- Gangliosidoses (e.g. Tay–Sachs disease: Tay’s sign);
- Metachromatic leucodystrophy;
- Niemann–Pick disease (especially type A).

Storage of sphingolipids or other substances in ganglion cells in the perimacular region gives rise to the appearance.

**Reference**

**Cross Reference**
Maculopathy

**Cheyne–Stokes Breathing**
- see PERIODIC RESPIRATION

**Chicken Wings**
In facioscapulohumeral (FSH) muscular dystrophy, the bulk of the deltoid and forearm muscles is normally well preserved, whilst biceps and triceps are wasted (and may be weak), thus giving rise to an appearance of the upper limbs sometimes labelled as ‘chicken wings’ or ‘Popeye arms’.

**Cross Reference**
Winging of the scapula

**Chorea, Choreaathetosis**
Chorea is an involuntary movement disorder characterized by jerky, restless, purposeless movements (literally dance-like) which tend to flit from one part of the body to another in a rather unpredictable way, giving rise to a fidgety appearance. There may also be athetoid movements (slow, sinuous, writhing), jointly referred to as choreoathetosis. Severe proximal choreiform movements of large amplitude (‘flinging’) are referred to as ballism or ballismus. When, as is often the case, such movements are confined to one side of the body they are referred to as hemichorea–hemiballismus. There may be concurrent abnormal muscle tone,
either hypotonia or rigidity. Hyperpronation of the upper extremity may be seen when attempting to maintain an extended posture.

The pathophysiology of chorea (as for ballismus) is unknown; movements may be associated with lesions of the contralateral subthalamic nucleus, caudate nucleus, putamen, and thalamus. One model of basal ganglia function suggests that reduced basal ganglia output to the thalamus disinhibits thalamic relay nuclei leading to increased excitability in thalamocortical pathways which passes to descending motor pathways resulting in involuntary movements. Recognized causes of chorea and choreoathetosis are many, including

- **Inherited disorders:**
  - **Autosomal dominant:**
    - Huntington’s disease;
    - Spinocerebellar ataxias: including SCA, Machado–Joseph disease, DRPLA;
    - Benign hereditary chorea (BHC).
  - **Autosomal recessive:**
    - Aminoacidopathies;
    - Ataxia telangiectasia (AT);
    - Basal ganglia calcification;
    - Lesch–Nyhan syndrome;
    - Lysosomal disorders;
    - Neuroacanthocytosis;
    - Neurodegeneration with brain iron accumulation (Hallervorden–Spatz disease);
    - Porphyria;
    - Tuberous sclerosis;
    - Urea cycle disorders;
    - Wilson’s disease.
  - **Others:**
    - Paroxysmal dyskinesias: paroxysmal kinesigenic choreoathetosis (PKC) and paroxysmal dystonic choreoathetosis (PDC);
    - Leigh’s syndrome;
    - Mitochondrial disease.

- **Drug-induced:**
  - Neuroleptics;
  - Propofol;
  - Antiepileptic drugs;
  - Antiparkinsonian medication: levodopa therapy in later stages of idiopathic Parkinson’s disease;
  - Oral contraceptives;
  - Amphetamines and tricyclic antidepressants (rare).

- **Toxic/metabolic:**
  - Alcohol;
  - Anoxia;
  - Carbon monoxide poisoning;
Cocaine;
Heavy metal poisoning;
Hyperthyroidism;
Hypoparathyroidism;
Pregnancy: chorea gravidarum.
Hypernatraemia or hyponatraemia, hypomagnesaemia, hypocalcaemia; hyperosmolality;
Hyperglycaemia or hypoglycaemia;
Non-Wilsonian acquired hepatocerebral degeneration;
Nutritional.

- **Infection:**
  - Sydenham’s chorea (postinfectious, rheumatic chorea, St Vitus dance, PANDAS);
  - Brainstem encephalitis, encephalitis lethargica;
  - Prion disease: Creutzfeldt–Jakob disease, variant CJD.

- **Immunological:**
  - Systemic lupus erythematosus;
  - Henoch–Schönlein purpura;
  - Neurosarcoidosis;
  - Multiple sclerosis;
  - Behçet’s disease (rare);
  - Vasculitis;
  - Hashimoto’s encephalopathy.

- **Vascular:**
  - Infarction (including Binswanger’s encephalopathy);
  - Haemorrhage;
  - Arteriovenous malformation;
  - Polycythaemia rubra vera (hyperviscosity);
  - Migraine;
  - Cerebral palsy.

- **Tumours:**
  - Primary and secondary (rare).

- **Others:**
  - Trauma;
  - Physiological chorea of infancy;
  - ‘Senile chorea’;
  - Postpump (cardiac bypass) chorea;
  - Psychogenic.

Where treatment is necessary, antidopaminergic agents such as dopamine-receptor antagonists (e.g. neuroleptics, sulpiride, risperidone) and dopamine-depleting agents (e.g. tetrabenazine, reserpine) may help, although they may cause parkinsonism, akathisia, neuroleptic malignant syndrome, and sedation. Chronic neuroleptic use may also cause chorea, but these movements are repetitive and predictable, unlike ‘classic’ chorea.
Chvostek's Sign

References

Cross References
Athetosis; Ballism, Ballismus; Dyskinesia; Hypotonia, Hypotonus; Milkmaid’s grip; Pseudochoreoathetosis; Rigidity; Trombone tongue

Chromaesthesia
- see SYNAESTHESIA

Chronognosia
This name has been sometimes given to a primary disturbance of the sense of time. Luria claimed it was associated with deep-seated temporal and temporo-diencephalic lesions, possibly right-sided lesions in particular. It occurs in some patients with Alzheimer’s disease who get up and dress, make tea, or phone relatives in the small hours, oblivious to the actual time, much to the exasperation of their loved ones. Whether this is a true agnosia remains open to investigation.

Reference

Cross Reference
Agnosia

Chvostek’s Sign
Chvostek’s sign is contraction of facial muscles provoked by lightly tapping over the facial nerve as it crosses the zygomatic arch. Chvostek’s sign is observed in hypocalcaemic states, such as hypoparathyroidism and the respiratory alkalosis associated with hyperventilation. There may be concurrent posturing of the hand, known as main d’accoucheur for its resemblance to the posture adopted for manual delivery of a baby.

The pathophysiology of this mechanosensitivity of nerve fibres is uncertain, but is probably related to increased discharges in central pathways. Although hypocalcaemia might be expected to impair neuromuscular junction transmission and excitation–contraction coupling (since Ca$^{2+}$ ions are required for these processes) this does not in fact occur.

Cross References
Main d’accoucheur; Spasm
**Ciliospinal Response**  
The ciliospinal response consists of rapid bilateral pupillary dilatation and palpebral elevation in response to a painful stimulus in the mantle area, for example, pinching the skin of the neck.  
**Reference**  
**Cross Reference**  
Pupillary reflexes

**Cinematic Vision**  
Cinematic vision is a form of metamorphopsia, characterized by distortion of movement with action appearing as a series of still frames as if from a movie. Causes include migraine aura, partial seizures, and schizophrenic psychosis.  
**Cross Reference**  
Metamorphopsia

**Circumlocution**  
Circumlocution may be used to refer to:  
- A discourse that wanders from the point, only eventually to return to the original subject matter, as seen in fluent aphasias.  
- A response to word-finding difficulties, as in early Alzheimer’s disease or non-fluent aphasias: in response to familiar pictures, patients may comment that the name is on the tip-of-the-tongue but they cannot access it, and therefore give alternatives, e.g. ‘gardener’s friend’ or ‘beetle’ for ladybird.  
**Reference**  
**Cross References**  
Anomia; Aphasia; Dementia

**Clapping Test**  
- see APPLAUSE SIGN

**Clasp-Knife Phenomenon**  
Clasp-knife phenomenon is the name sometimes applied to the sudden ‘give’ encountered when passively moving a markedly spastic limb. Since the clasp-knife phenomenon is a feature of spasticity, the term ‘clasp-knife rigidity’ is probably best eschewed to avoid possible confusion.  
**Cross References**  
Rigidity; Spasticity

**Claudication**  
Claudication (literally limping, Latin claudicatio) refers to intermittent symptoms of pain secondary to ischaemia. Claudication of the legs on walking is a symptom of peripheral vascular disease. Claudication of the jaw, tongue, and limbs (especially upper) may be a feature of giant cell (temporal) arteritis. Jaw
Clonus is said to occur in 40% of patients with giant cell arteritis and is the presenting complaint in 4%; tongue claudication occurs in 4% and is rarely the presenting feature. Presence of jaw claudication is one of the clinical features which increases the likelihood of a positive temporal artery biopsy.

**Reference**

**Claw Foot**
Claw foot, or *pied en griffe*, is an abnormal posture of the foot, occurring when weakness and atrophy of the intrinsic foot muscles allows the long flexors and extensors to act unopposed, producing shortening of the foot, heightening of the arch, flexion of the distal phalanges and dorsiflexion of the proximal phalanges (cf. pes cavus). This may occur in chronic neuropathies of early onset which involve motor fibres, such as hereditary motor and sensory neuropathies (types I and II).

**Cross Reference**
Pes cavus

**Claw Hand**
Claw hand, or *main en griffe*, is an abnormal posture of the hand with hyperextension at the metacarpophalangeal joints (fifth, fourth, and, to a lesser extent, third finger) and flexion at the interphalangeal joints. This results from ulnar nerve lesions above the elbow, or injury to the lower part of the brachial plexus (Dejerine–Klumpke type), producing wasting and weakness of hypothenar muscles, interossei, and ulnar (medial) lumbricals, allowing the long finger extensors and flexors to act unopposed.

**Reference**

**Cross References**
Benediction hand; Camptodactyly

**Clonus**
Clonus is rhythmic, involuntary, repetitive, muscular contraction and relaxation. It may be induced by sudden passive stretching of a muscle or tendon, most usually the Achilles tendon (ankle clonus) or patella (patellar clonus). Ankle clonus is best elicited by holding the relaxed leg underneath the moderately flexed knee, then quickly dorsiflexing the ankle and holding it dorsiflexed. A few beats of clonus are within normal limits but sustained clonus is pathological.

Clonus reflects hyperactivity of muscle stretch reflexes and may result from self-re-excitation. It is a feature of upper motor neurone disorders affecting the corticospinal (pyramidal) system. Patients with disease of the corticospinal tracts may describe clonus as a rhythmic jerking of the foot, for example, when using the foot pedals of a car. Clonus may also be observed as part of a generalized (primary or secondary) epileptic seizure, either in isolation (clonic seizure) or much more commonly following a tonic phase (tonic–clonic seizure). The clonic movements usually involve all four limbs and decrease in frequency and increase
in amplitude over about 30–60 s as the attack progresses. Rather different ‘clonic’ movements may occur in non-epileptic seizures. A few clonic jerks may also be observed in syncopal attacks, leading the uninitiated to diagnose ‘seizure’ or ‘convulsion’.

**Cross References**
Myoclonus; Seizure; Upper motor neurone (UMN) syndrome

**Closed Fist Sign**
This is one of the provocative tests for carpal tunnel syndrome: it is positive if paraesthesia in the distribution of the median nerve develops after maintaining fist closure for 60 s.

**References**

**Cross References**
Flick sign; Phalen’s sign; Tinel’s sign

**Closing-In Sign**
Copying of drawings which are close to or superimposed on the original has been referred to as the ‘closing-in’ sign. It may be seen in patients with Alzheimer’s disease with deficits in visuospatial function. This has sometimes been characterized as one aspect of the ‘constructional apraxia’ of Alzheimer’s disease; it may be useful in differentiating AD from subcortical vascular dementia.

**References**

**Cluster Breathing**
Damage at the pontomedullary junction may result in a breathing pattern characterized by a cluster of breaths following one another in an irregular sequence. This sign may be of localizing value in comatose patients.

**Cross Reference**
Coma

**Coactivation Sign**
This sign is said to be characteristic of psychogenic tremors, namely, increased tremor amplitude with loading (cf. reduced amplitude of organic tremor with loading), perhaps due to muscle coactivation to maintain oscillation.

**Reference**
Collapsing Weakness

Cross Reference
Tremor

Cock Walking
- see TOE WALKING

Cogan’s (Lid Twitch) Sign
Cogan’s sign is a twitching of the upper eyelid seen a moment after the eyes are moved from downgaze to the primary position. Twitches may also be seen with eye closure after sustained upgaze. These phenomena are said to be characteristic signs of ocular myasthenia gravis and were found in 60% of myasthenics in one study. They may also occur occasionally in other oculomotor brainstem disorders such as Miller Fisher syndrome, but are not seen in normals. Hence, the sign is neither sensitive nor specific.

Cogan’s lid twitch sign should not be confused with either Cogan’s syndrome, an autoimmune disorder of episodic vertigo, tinnitus, hearing loss, and interstitial keratitis; or the oculomotor apraxia of Cogan, a congenital lack of lateral gaze.

References

Cross References
Fatigue; Ice pack test; Ocular apraxia

Cogwheeling, Cogwheel Phenomenon, Cogwheel Rigidity
- see RIGIDITY; SACCADIC INTRUSION, SACCADIC PURSUIT

Cold Hands Sign
In multiple system atrophy (MSA), the hands may be cold, dusky, and violaceous with poor circulatory return after blanching by pressure, suggesting defective neurovascular control of the distal extremities as one feature of the autonomic dysfunction in MSA. The findings are not present in idiopathic Parkinson’s disease.

Reference

Collapsing Weakness
Collapsing weakness, or ‘give-way’ weakness, suggesting intermittent voluntary effort, is often taken as a sign of functional weakness. Although sometimes labelled as ‘volitional weakness’, it is not clear that such weakness is in any conscious sense willed, and it is therefore probably better to use a non-committal
term such as ‘apparent weakness’. Such collapsing weakness has also been recorded following acute brain lesions such as stroke.

References

Cross References
Functional weakness and sensory disturbance; Spasticity; Weakness; ‘Wrestler’s sign’

**Collier’s Sign**

Collier’s sign (‘posterior fossa stare’, ‘tucked lid’ sign), first described in 1927, is elevation and retraction of the upper eyelids, baring the sclera above the cornea, with the eyes in the primary position or looking upward. This may be seen with upper dorsal midbrain supranuclear lesions, e.g. ‘top of the basilar syndrome’, Parinaud’s syndrome. There may be accompanying paralysis of vertical gaze (especially upgaze) and light-near pupillary dissociation. The sign is thought to reflect damage to the posterior commissure levator inhibitory fibres.

References

Cross References
Lid retraction; Light-near pupillary dissociation; Parinaud’s syndrome

**Colour Anomia**

- see ACHROMATOPSIA; ANOMIA

**Coma**

Coma is a state of unresponsiveness, with eyes closed, from which a patient cannot be roused by verbal or mechanical stimuli. It represents a greater degree of impairment of consciousness than stupor or obtundation, all three forming part of a continuum, rather than discrete stages, ranging from alert to comatose. This lack of precision prompts some authorities to prefer the description of the individual aspects of neurological function in unconscious patients, such as eye movements, limb movements, vocalization, and response to stimuli, since this conveys more information than the use of terms such as coma, stupor, or obtundation, or the use of a lumped ‘score’, such as the Glasgow Coma Scale. These signs should be documented serially to assess any progression of coma. Assessment of the depth of coma may be made by observing changes in eye movements and response to central noxious stimuli: roving eye movements are lost before oculocephalic responses; caloric responses are lost to go. The switch from flexor to extensor posturing (decorticate vs. decerebrate rigidity) also indicates increasing depth of coma.
There are many causes of coma, which may be broadly categorized as structural or toxic-metabolic; the latter are generally more slowly progressive and produce symmetrical signs, whereas structural lesions more often have an abrupt onset and some focal asymmetric findings on examination, but these distinctions are not absolute. Recognized causes of coma include

- **Structural:**
  - Vascular insults (subarachnoid haemorrhage, cerebral infarction or haemorrhage, CADASIL);
  - Trauma;
  - Tumour;
  - Hydrocephalus;
  - Vasculitides, leucodystrophies, leucoencephalopathies.

- **Toxic-metabolic:**
  - Metabolic causes: e.g. hypoxia, hypercapnia, hypoglycaemia;
  - Infections: e.g. meningitis, encephalitis, sepsis;
  - Epilepsy.

Unrousability which results from psychiatric disease, or which is being feigned (‘pseudocoma’), also needs to be differentiated.

A number of neurobehavioural states may be mistaken for coma, including abulia, akinetic mutism, catatonia, and the locked-in syndrome. EEG features may assist in differential diagnosis: prominent rhythmic beta activity raises the possibility of drug intoxication.

**References**


**Cross References**

Abulia; Akinetic mutism; Caloric testing; Catatonia; Decerebrate rigidity; Decorticate rigidity; Locked-in syndrome; Obtundation; Oculocephalic response; Roving eye movements; Stupor; Vegetative states; Vestibulo-ocular reflexes

**Compulsive Grasping Hand**

This name has been given to involuntary left-hand grasping related to all right-hand movements in a patient with a callosal haemorrhage. This has been interpreted as a motor grasp response to contralateral hand movements and a variant of anarchic or alien hand. The description does seem to differ from that of behaviours labelled as forced groping and the alien grasp reflex.

**Reference**

Conduction Aphasia

Conduction aphasia is defined as a fluent aphasia with paraphasic errors (especially phonemic/literal) during speech, repetition, and naming. In its ‘pure’ form, there is a dissociation between relatively preserved auditory and reading comprehension of language and impaired repetition (in which the phenomenon of conduit d’approche may occur) and naming. Reading comprehension is good or normal and is better than reading aloud which is impaired by paraphasic errors.

Conduction aphasia was traditionally explained as due to a disconnection between sensory (Wernicke) and motor (Broca) areas for language, involving the arcuate fasciculus in the supramarginal gyrus. Certainly the brain damage (usually infarction) associated with conduction aphasia most commonly involves the left parietal lobe (lower postcentral and supramarginal gyri) and the insula, but it is variable, and the cortical injury may be responsible for the clinical picture.

Conduction aphasia is most often seen during recovery from Wernicke’s aphasia, and clinically there is often evidence of some impairment of comprehension. If isolated, the prognosis for conduction aphasia is good.

References

Conduit d’approche

Conduit d’approche, or ‘homing-in’ behaviour, is a verbal output phenomenon applied to patients with conduction aphasia attempting to repeat a target word, in which multiple phonemic approximations of the word are presented, with gradual improvement until the target word is achieved. This phenomenon suggests that an acoustic image of the target word is preserved in this condition. A similar phenomenon may be observed in patients with optic aphasia attempting to name a visual stimulus. A similar behaviour is seen in so-called speech apraxia, in which patients repeatedly approximate to the desired output before reaching it.

The term may also be used to refer to a parapraxis in which patients attempt to perform a movement several times before achieving the correct movement.

Cross References
Aphasia; Conduction aphasia; Optic aphasia; Parapraxia, Parapraxis; Speech apraxia
**Confabulation**
The old definition of confabulation as the falsification of episodic memory occurring in clear consciousness, often in association with amnesia (in other words, paramnesias related as true events), has proven increasingly deficient, not least because most amnesic patients, suffering from medial temporal lobe/hippocampal lesions, do not confabulate, and poor memory alone cannot explain confabulation. Schnider has developed a fourfold schema of intrusions, momentary confabulations, fantastic confabulations, and behaviourally spontaneous confabulations, of which the latter are clinically the most challenging. Anterior limbic structures are thought culpable, and the pathogenesis includes a wide variety of diseases, which may include associated phenomena such as amnesia, disorientation, false recognition syndromes including the Capgras delusion, and anosognosia. Psychophysical and neuroimaging studies suggest that confabulators have reality confusion and a failure to integrate contradictory information due to the failure of a filtering process, 200–300 ms after stimulus presentation and before recognition and re-encoding, which normally permits suppression of currently irrelevant memories.

**References**

**Cross References**
Amnesia; Asomatognosia; Cortical blindness; Delusion; Paramnesia

**Confusion**
Confusion, understood as the inability to think with one’s customary clarity and coherence, is a feature of not only delirium, but also of other situations (encephalopathies, attentional disorders). Moreover, as there is a lack of correlation of meaning when this term is used by different health professionals, it is regarded by some as an unhelpful term.

**References**

**Cross Reference**
Delirium

**Congenital Nystagmus**
Congenital nystagmus is a pendular nystagmus with the following characteristics:

- Usually noted at birth or in early infancy; sometimes may only become apparent in adult life;
- Irregular waveforms;
- Conjugate;
- Almost always horizontal;
Consensual Light Reflex

- Accentuated by fixation, attention, anxiety;
- Decreased by convergence, active eyelid closure;
- Often a null point or region;
- No complaint of oscillopsia;
- It may appear with blindness of childhood onset.

Acquired pendular nystagmus may be a result of neurological disease which may present in childhood, such as Pelizaeus–Merzbacher disease, mitochondrial disease, multiple sclerosis, and Whipple’s disease.

Cross References
Nystagmus; Oscillopsia

Consensual Light Reflex
- see PUPILLARY REFLEXES

Constructional Apraxia
- see APRAXIA

Contracture

The term contracture may be used in various contexts:

- Clinically, to describe an acquired restriction of joint mobility (prenatally acquired restriction of joint mobility is called arthrogryposis). This may be due to a variety of factors, including prolonged muscle spasticity with or without muscle fibrosis (i.e. without pathological muscle shortening) and ligamentous restrictions. This often occurs in the context of limb immobilization or inactivity, for example, in a flexed posture. Injections of botulinum toxin to abolish muscle spasticity may be required to assess whether there is concurrent ligamentous restriction, and thus to plan optimum treatment, which may involve surgery. Contractures of muscular origin may be seen in conditions such as Emery–Dreifuss disease (especially elbow, Achilles tendon, posterior part of neck), Bethlem myopathy, and Ullrich congenital muscular dystrophy associated with mutations in genes encoding the peptide chains of collagen VI, limb girdle muscular dystrophy type 2A associated with mutations in the calpain 3 gene, and Duchenne muscular dystrophy.
- Clinically, to describe a hard, contracted muscle which is painful to straighten and lasting for several hours following exercise in a metabolic myopathy such as McArdle’s disease (myophosphorylase deficiency, glycogen storage disease type V); this may be associated with EMG silence.
- Physiologically, to describe a prolonged painful muscle spasm with EMG silence, as observed in myotonia and paramyotonia.

Reference

Cross References
Myotonia; Paramyotonia; Paraplegia; Spasm; Spasticity
**Corneal Reflex**

**Convergence–Retraction Nystagmus**
- see NYSTAGMUS; PARINAUD’S SYNDROME

**Coprolalia**
Coprolalia is the use of expletives or other obscene language. This may be
- **Vocal**: involuntary utterance of obscenities;
- **Mental**: compulsion to think obscenities.

The former is a complex vocal tic most characteristically seen in Tourette syndrome although it actually occurs in less than half of affected individuals. Other recognized disease associations are as follows:
- Lesch–Nyhan syndrome;
- Postencephalitic parkinsonism;
- Neuroacanthocytosis;
- Cingulate cortical seizures.

The pathophysiology of coprolalia is unknown but may be related to frontal (cingulate and orbitofrontal) dysfunction, for which there is some evidence in Tourette syndrome.

**Reference**

**Cross Reference**
Tic

**Copropraxia**
Copropraxia is a complex motor tic comprising obscene gesturing, sometimes seen in Tourette syndrome.

**Cross References**
Coprolalia; Tic

**Corectopia**
Corectopia is pupillary displacement, which may be seen with midbrain lesions, including transtentorial herniation and top-of-the-basilar syndrome, peripheral oculomotor nerve palsies, and focal pathology in the iris.

**Reference**

**Corneal Reflex**
The corneal reflex consists of a bilateral blink response elicited by touching the cornea lightly, for example, with a piece of cotton wool. As well as observing whether the patient blinks, the examiner should also ask whether the stimulus was felt: a difference in corneal sensitivity may be the earliest abnormality in this reflex. Synkinetic jaw movement may also be observed: the corneomandibular reflex.

The afferent limb of the corneal reflex is via the trigeminal (V) nerve, the efferent limb via the facial (VII) nerve to orbicularis oculi. The fibres subserving
the corneal reflex seem to be the most sensitive to trigeminal nerve compression or distortion: an intact corneal reflex with a complaint of facial numbness leads to suspicion of a non-organic cause. Reflex impairment may be an early sign of a cerebellopontine angle lesion, which may also cause ipsilateral lower motor neurone type facial (VII) weakness and ipsilateral sensorineural hearing impairment (VIII). Trigeminal nerve lesions cause both ipsilateral and contralateral corneal reflex loss.

Cerebral hemisphere (but not thalamic) lesions causing hemiparesis and hemisensory loss may also be associated with a decreased corneal reflex.

The corneal reflex has a high threshold in comatose patients and is usually preserved until late (unless coma is due to drug overdose), in which case its loss is a poor prognostic sign.

**Cross References**
Blink reflex; Coma; Cerebellopontine angle syndrome; Corneomandibular reflex; Facial paresis

**Corneomandibular Reflex**
The corneomandibular reflex, also known as the corneopterygoid reflex or Wartenberg’s reflex or sign, consists of anterolateral jaw movement following corneal stimulation. In one study, the corneomandibular reflex was observed in about three-quarters of patients with motor neurone disease (MND) who displayed no other pathological reflexes, a frequency much higher than that seen in patients with stroke causing hemiparesis or pseudobulbar palsy. It was therefore suggested to be a sensitive indicator of upper motor neurone involvement in MND.

**References**

**Cross References**
Corneal reflex; Pseudobulbar palsy

**Corneopterygoid Reflex**
- see CORNEOMANDIBULAR REFLEX

**Cortical Blindness**
Cortical blindness (*Rindblindheit*) is loss of vision due to bilateral visual cortical damage, usually hypoxic–ischaemic in origin, or bilateral subcortical lesions affecting the optic radiations. A small central field around the fixation point may be spared (macula sparing). Pupillary reflexes are preserved but optokinetic nystagmus cannot be elicited.

Cortical blindness may result from:
- Bilateral (sequential or simultaneous) posterior cerebral artery occlusion;
- ‘Top of the basilar syndrome’;
- Migraine;
- Cerebral anoxia;
Coup de Sabre

- Bacterial endocarditis;
- Wegener’s granulomatosis;
- Following coronary or cerebral angiography (may be transient);
- Epilepsy (transient);
- Ciclosporin therapy, e.g. following organ transplantation.

If acute in onset (i.e. vascular), cortical blindness may ultimately evolve to prosopagnosia via visual object agnosia.

Patients with cortical blindness may deny their visual defect (Anton’s syndrome, visual anosognosia) and may confabulate about what they ‘see’.

Cross References
Anosognosia; Confabulation; Macula sparing, Macula splitting; Optokinetic nystagmus, Optokinetic response; Prosopagnosia; Pupillary reflexes; Visual agnosia

Cotard’s Syndrome
A delusional syndrome, first described in the 1890s, characterized by the patient’s denial of their own existence, or of part of their body. The patient may assert that they are dead and able to smell rotten flesh or feel worms crawling over their skin. Although this may occur in the context of psychiatric disease, especially depression and schizophrenia, it may also occur in association with organic brain abnormalities, specifically lesions of the non-dominant temporoparietal cortex, or migraine.

Some envisage Cotard’s syndrome as a more pervasive form of the Capgras syndrome, originating similarly as a consequence of Geschwindian disconnection between the limbic system and all sensory areas, leading to a loss of emotional contact with the world. Antidepressant treatment and/or ECT may sometimes be helpful in Cotard syndrome of psychiatric origin.

References

Cross References
Capgras syndrome; Delusion; Disconnection syndromes

Coup de Poignard
*Coup de poignard*, or dagger thrust, refers to a sudden precordial pain, as may occur in myocardial infarction or aortic dissection, also described with spinal subarachnoid haemorrhage.

References

Coup de Sabre
*Coup de sabre* is a localized form of scleroderma manifest as a linear, atrophic lesion on the forehead which may be mistaken for a scar. This lesion may be associated with hemifacial atrophy and epilepsy, and neuroimaging may
show hemiatrophy and intracranial calcification. Whether these changes reflect inflammation or a neurocutaneous syndrome is not known.

Reference

Cross Reference
Hemifacial atrophy

**Cover Tests**
The simple cover and cover–uncover tests may be used to demonstrate manifest and latent strabismus (heterotropia and heterophoria), respectively.

The cover test demonstrates tropias: the uncovered eye is forced to adopt fixation; any movement therefore represents a manifest strabismus (heterotropia).

The cover–uncover test demonstrates phorias: any movement of the covered eye to re-establish fixation as it is uncovered represents a latent strabismus (heterophoria).

The alternate cover or cross-cover test, in which the hand or occluder moves back and forth between the eyes, repeatedly breaking and re-establishing fixation, is more dissociating, preventing binocular viewing, and therefore helpful in demonstrating whether or not there is strabismus. It should be performed in the nine cardinal positions of gaze to determine the direction that elicits maximal deviation. However, it does not distinguish between tropias and phorias, for which the cover and cover–uncover tests are required.

Cross References
Heterophoria; Heterotropia

**Cramp**
Cramps are defined as involuntary contractions of a number of muscle units which results in a hardening of the muscle with pain due to a local lactic acidosis. Cramps are not uncommon in normal individuals but in a minority of cases they are associated with an underlying neurological or metabolic disorder. Cramps need to be distinguished from spasticity, neuromyotonia, and myokymia. Recognized associations of cramp include

- **Normal individuals:**
  - Especially during periods of dehydration with salt loss; pregnancy.
  - Benign cramp syndrome, there is a family history of cramps.

- **Metabolic causes:**
  - Hypothyroidism;
  - Haemodialysis;
  - Hypocalcaemia; hyperventilation (with secondary hypocalcaemia).

- **Neurological causes:**
  - Chronic peripheral neuropathy;
  - Metabolic myopathies (e.g. myophosphorylase deficiency, lactate dehydrogenase (LDH) deficiency, with exercise intolerance and myoglobinuria);
  - Muscular dystrophies (especially Becker, Duchenne);
Crossed Adductor Reflex

Motor neurone disease;
Stiff man syndrome.

Treatment involves addressing any underlying metabolic abnormality. Symptomatic treatment of cramps may include use of quinine sulphate, vitamin B, naftidrofuryl, and calcium channel antagonists such as diltiazem; carbamazepine, phenytoin, and procainamide have also been tried.

References

Cross References
Fasciculation; Myokymia; Neuromyotonia; Spasm; Stiffness

Cremasteric Reflex
The cremasteric reflex is a superficial or cutaneous reflex consisting of contraction of the cremaster muscle causing elevation of the testicle, following stimulation of the skin of the upper inner aspect of the thigh from above downwards (i.e. the L1, L2 dermatomes, via the ilioinguinal and genitofemoral nerves).

The cremasteric reflex is lost when the corticospinal pathways are damaged above T12 or following lesions of the genitofemoral nerve. It may also be absent in elderly men or with local pathology such as hydrocele, varicocele, orchitis, or epididymitis.

Cross References
Abdominal reflexes; Reflexes

Crocodile Tears
Crocodile tears, gustatory epiphora, or Bogorad’s syndrome reflect inappropriate unilateral lacrimation during eating, such that tears may spill down the face (epiphora). This autonomic synkinesis is a striking but rare consequence of aberrant reinnervation of the facial (VII) nerve, usually after a Bell’s palsy, when fibres originally supplying the salivary glands are re-routed to the lacrimal gland via the greater superficial petrosal nerve.

Cross References
Bell’s palsy; Epiphora; Synkinesia, Synkinesis

Crossed Adductor Reflex
Contralateral adductor muscle contraction in response to a tap on the adductor tendon may be found with a pyramidal lesion above L2, although it is a normal finding in infants.

Cross Reference
Reflexes
Crossed Aphasia
Aphasia from a right-sided lesion in a right-handed patient, crossed aphasia, is rare, presumably a reflection of crossed or mixed cerebral dominance. It may occur transiently during a focal epileptic seizure or migraine aura.

References

Cross Reference
Aphasia

Crossed Apraxia
A name given to apraxia in right-handed patients with right-sided lesions; apraxia is more commonly associated with left-sided brain injury.

Reference

Cross Reference
Apraxia

Crossed Straight Leg Raising
- see LASÈGUE’S SIGN

Cross over
In the line bisection task for the detection of unilateral spatial neglect, in which the subjective midline is placed more towards the ipsilesional extreme of the line compared to the objective midline, especially with longer lines (length effect), with shorter lines there is a paradoxical deviation towards the contralesional side, a sign called cross-over.

Reference

Cross Reference
Neglect

Crying
- see AUTOMATISM; PATHOLOGICAL CRYING, PATHOLOGICAL LAUGHTER; SEIZURES

Cuirasse
- see SUSPENDED SENSORY LOSS

Cushing Reflex, Cushing Response
This is the triad of increasing systolic and pulse pressure with bradycardia and slow irregular respiration associated with increased intracranial pressure which may lead to cerebral herniation and fatal brainstem compression, for example, with posterior fossa masses or subarachnoid haemorrhage.
Reference

Czarnecki’s Sign
Aberrant regeneration of the oculomotor (III) nerve to the iris sphincter may lead to gaze-evoked segmental constriction of the pupil, which may be visible only with slit-lamp examination.

Reference
Dalrymple’s Sign
Dalrymple’s sign is increased width of the palpebral fissure, often seen in hyperthyroidism.

Cross Reference
Lid retraction

Dazzle
Dazzle is a painless intolerance of the eyes to bright light (cf. photophobia). It may be peripheral in origin (retinal disease; opacities within cornea, lens, vitreous); or central (lesions anywhere from optic nerve to occipitotemporal region).

Cross Reference
Photophobia

Decerebrate Rigidity
Decerebrate rigidity is a posture observed in comatose patients in which there is extension and pronation of the upper extremities, extension of the legs, and plantar flexion of the feet (= extensor posturing), which is taken to be an exaggeration of the normal standing position. Painful stimuli may induce opisthotonos, hyperextension, and hyperpronation of the upper limbs.

Decerebrate rigidity occurs in severe metabolic disorders of the upper brainstem (anoxia/ischaemia, trauma, structural lesions, drug intoxication). A similar picture was first observed by Sherrington (1898) following section of the brainstem of cats at the collicular level, below the red nuclei, such that the vestibular nuclei were intact. The action of the vestibular nuclei, unchecked by higher centres, may be responsible for the profound extensor tone.

Decerebrate rigidity indicates a deeper level of coma than decorticate rigidity; the transition from the latter to the former is associated with a worsening of prognosis.

Reference

Cross References
Coma; Decorticate rigidity; Opisthotonos

De Clérambault Syndrome
- see DELUSION

Decomposition of Movement
- see ASYNERGIA
Decorticate Rigidity
Decorticate rigidity is a posture observed in comatose patients in which there is adduction of the shoulders and arms and flexion of the elbows and wrists (= flexor posturing). The lesion responsible for decorticate rigidity is higher in the neuraxis than that causing decerebrate rigidity, often being diffuse cerebral hemisphere or diencephalic disease, although, despite the name, it may occur with upper brainstem lesions. Common causes are anoxia/ischaemia, trauma, and drugs.
Cross References
Coma; Decerebrate rigidity

Déjà Entendu
A sensation of familiarity akin to déjà vu but referring to auditory rather than visual experiences.

Déjà Vécu
- see Déjà VU

Déjà Vu
Déjà vu (literally ‘already seen’) is a subjective, inappropriate impression of familiarity for a present experience in relation to an undefined past. However, since the term has passed into the vernacular, not every patient complaining of ‘déjà vu’ has a pathological problem. The term may be used colloquially to indicate familiar events or experiences. Recurrent hallucinations or vivid dream-like imagery may also enter the differential diagnosis. A phenomenon of slight confusion in which all is not clear although it is familiar has sometimes been labelled ‘prèsque vu’.

Epileptic déjà vu may last longer and be more frequent and may be associated with other features such as depersonalization and derealization, strong emotion such as fear, epigastric aura, or olfactory hallucinations. Epileptic déjà vu is a complex aura of focal onset epilepsy; specifically, it is indicative of temporal lobe onset of seizures and is said by some authors to be the only epileptic aura of reliable lateralizing significance (right). Déjà vécu (‘already lived’) has been used to denote a broader experience than déjà vu but the clinical implications are similar.

Déjà vu has also been reported to occur in several psychiatric disorders, such as anxiety, depression, and schizophrenia.

References

Cross References
Aura; Hallucination; Jamais vu

Delirium
Delirium, also sometimes known as acute confusional state, acute organic reaction, acute brain syndrome, or toxic-metabolic encephalopathy, is a neurobehavioural syndrome of which the cardinal feature is a deficit of attention, the ability to focus on specific stimuli. Diagnostic criteria also require a concurrent
Delirium

alteration in level of awareness, which may range from lethargy to hypervigilance, although delirium is not primarily a disorder of arousal or alertness (cf. coma, stupor, obtundation). Other features commonly observed in delirium include

- **impaired cognitive function**: disorientation in time and place;
- **perceptual disorders**: illusions, hallucinations;
- **behavioural disturbances**: agitation, restlessness, aggression, wandering, which may occur as a consequence of perceptual problems (hyperalert type); or unresponsiveness, withdrawal (hypoalert or quiet type);
- **language**: rambling incoherent speech, logorrhoea;
- **altered sleep–wake cycle**: ‘sundowning’ (restlessness and confusion at night);
- tendency to marked fluctuations in alertness/activity, with occasional lucid intervals;
- **delusions**: often persecutory.

Hence this abnormal mental state shows considerable clinical heterogeneity. Subtypes or variants are described, one characterized by hyperactivity (‘agitated’), the other by withdrawal and apathy (‘quiet’).

The course of delirium is usually brief (seldom more than a few days, often only hours). On recovery the patient may have no recollection of events, although islands of recall may be preserved, corresponding with lucid intervals (a useful, if retrospective, diagnostic feature).

Delirium is often contrasted with dementia, a ‘chronic brain syndrome’, in which attention is relatively preserved, the onset is insidious rather than acute, the course is stable over the day rather than fluctuating, and which generally lasts months to years. However, it should be noted that in the elderly delirium is often superimposed on dementia, which is a predisposing factor for the development of delirium, perhaps reflecting impaired cerebral reserve.

The pathophysiology of delirium is not well understood. Risk factors for the development of delirium may be categorized as either predisposing or precipitating.

- **Predisposing factors** include
  - Age: frailty, physiological age rather than chronological
  - Sex: men > women
  - Neurological illness: dementia
  - Burden of comorbidity; dehydration
  - Drugs: especially anticholinergic medication
  - Primary sensory impairment (hearing, vision)

- **Precipitating factors** include
  - Drugs/toxins: benzodiazepines, opiates
  - Alcohol, especially withdrawal from, as in delirium tremens
  - Intercurrent illness:
    - Infection: primary CNS (encephalitis, meningitis), or systemic (urinary tract, chest, sepsis)
    - Metabolic: hypoxia, hypo-/hyperglycaemia, hepatic failure, uraemia, porphyria
CNS disorders: head injury, cerebrovascular disease, epilepsy (e.g. some forms of status), inflammatory disorders (e.g. collagen vascular disease)

Iatrogenic events: surgery (especially cardiac, orthopaedic)

These precipitating factors merit treatment in their own right, and investigations should be tailored to identify these aetiological factors. The EEG may show non-specific slowing in delirium, the degree of which is said to correlate with the degree of impairment, and reverses with resolution of delirium.

It is suggested that optimal nursing of delirious patients should aim at environmental modulation to avoid both understimulation and overstimulation; a side room is probably best (if possible). Drug treatment is not mandatory, the evidence base for pharmacotherapy is slim. However, if the patient poses a risk to him/herself, other patients, or staff which cannot be addressed by other means, regular low-dose oral haloperidol may be used, probably in preference to atypical neuroleptics, benzodiazepines (lorazepam), or cholinesterase inhibitors.

References

Cross References
Agraphia; Attention; Coma; Delusion; Dementia; Hallucination; Illusion; Logorrhoea; Obtundation; Stupor; ‘Sundowning’

Delusion
A delusion is a fixed false belief, not amenable to reason (i.e. held despite evidence to the contrary), and not culturally sanctioned. There are a number of common forms of delusion, including

• persecutory (paranoia);
• reference: important events or people being influenced by patient’s thoughts, ideas;
• grandiose/expansive: occur particularly in mania;
• guilt/worthlessness: occur particularly in depression;
• hypochondria;
• thought broadcast and thought insertion;
• control by an external agency.

Specific, named, delusional syndromes are those of:

• Capgras: the ‘delusion of doubles’, a familiar person or place is thought to be an impostor, or double; this resembles the reduplicative paramnesia described in neurological disorders such as Alzheimer’s disease.
• Fregoli: a familiar person is identified in other people, even though they bear no resemblance; this may occur in schizophrenia.
Dementia

- **De Clérambault** (erotomania): the belief (usually of a single woman) that a famous person is secretly in love with her (‘hope’), followed by the belief that the person is persecuting her (‘resentment’); may occur in schizophrenia.

Delusions are not only a feature of primary psychiatric disease (psychoses such as schizophrenia; neuroses such as depression), but may also be encountered in neurological disease with secondary psychiatric features (‘organic psychiatry’), e.g. delirium, and dementing syndromes such as Alzheimer's disease, dementia with Lewy bodies.

**Reference**

**Cross References**
Delirium; Dementia; Hallucination; Illusion; Intermetamorphosis; Misidentification syndromes; Reduplicative paramnesia

**Dementia**

Dementia is a syndrome characterized by loss of intellectual (cognitive) functions sufficient to interfere with social and occupational functioning. Cognition encompasses multiple functions including language, memory, perception, praxis, attentional mechanisms, and executive function (planning, reasoning). These elements may be affected selectively or globally: older definitions of dementia requiring global cognitive decline have now been superseded. Amnesia may or may not, depending on the classification system used, be a sine qua non for the diagnosis of dementia. Attentional mechanisms are largely preserved, certainly in comparison with delirium, a condition which precludes meaningful neuropsychological assessment because of profound attentional deficits. Multiple neuropsychological tests are available to test different areas of cognition.

Although more common in the elderly, dementia can also occur in the pre-senium and in children who may lose cognitive skills as a result of hereditary metabolic disorders. Failure to develop cognitive skills is termed learning disability. The heterogeneity of dementia is further exemplified by the fact that it may be acute or insidious in onset, and its course may be progressive, stable, or, in some instances, reversible (‘dysmentia’). A distinction is drawn by some authors between cortical and subcortical dementia: in the former the pathology is predominantly cortical and neuropsychological findings are characterized by amnesia, agnosia, apraxia, and aphasia (e.g. Alzheimer’s disease); in the latter pathology is predominantly frontal–subcortical and neuropsychological deficits include psychomotor retardation, attentional deficits, with relative preservation of memory and language; movement disorders may also be apparent (e.g. progressive supranuclear palsy, Huntington’s disease). However, not all authors subscribe to this distinction and considerable overlap may be observed clinically.

Cognitive deficits also occur in affective disorders such as depression, usually as a consequence of impaired attentional mechanisms. This syndrome is often labelled as ‘pseudodementia’ since it is potentially reversible with treatment of the underlying affective disorder. It may be difficult to differentiate dementia originating from depressive or neurodegenerative disease, since depression may also
Dementia

be a feature of the latter. Impaired attentional mechanisms may account for the common complaint of not recalling conversations or instructions immediately after they happen (aprosexia). Behavioural abnormalities are common in dementias due to degenerative brain disease and may require treatment in their own right.

Recognized causes of a dementia syndrome include

- **Neurodegenerative diseases:**
  - Alzheimer’s disease, frontaltemporal lobar degenerations (frontotemporal dementia, encompassing Pick’s disease; semantic dementia; primary non-fluent aphasia), dementia with Lewy bodies, Huntington’s disease, progressive supranuclear palsy, corticobasal degeneration, prion disease, Down’s syndrome, dementia pugilistica.

- **Cerebrovascular disease:**
  - Focal strategic infarcts (e.g. paramedian thalamic infarction), multiple infarcts, subcortical vascular disease,Binswanger’s disease.

- **Inflammatory disorders:** multiple sclerosis, systemic lupus erythematosus.

- **Structural disease:** normal pressure hydrocephalus, subdural haematoma, tumours, dural arteriovenous fistula.

- **Infection:** HIV dementia, neurosyphilis, Whipple’s disease.

- **Metabolic causes:** Wernicke–Korsakoff syndrome, vitamin B$_{12}$ deficiency, hypothyroidism, hyperparathyroidism/hypercalcaemia, leukodystrophies, Wilson’s disease.

Cognitive dysfunction may be identified in many other neurological illnesses. Investigation of patients with dementia aims to identify its particular cause. Because of the possibility of progression, reversible causes are regularly sought though very rare. Specific treatments for dementia are few: cholinesterase inhibitors have been licensed for the treatment of mild-to-moderate Alzheimer’s disease and may find a role in other conditions, such as dementia with Lewy bodies and vascular dementia, for behavioural as well as mnemonic features.

**References**


**Cross References**

Agnosia; Amnesia; Aphasia; Apraxia; Aprosexia; Attention; Delirium; Dysmentia; Pseudodementia; Psychomotor retardation
Depersonalization
Depersonalization, a form of dissociation, is the experience of feeling detached or alienated from oneself, such that the body feels strange, lacking control, or being viewed from the outside. There may be concurrent derealization. Depersonalization is a very common symptom in the general population and may contribute to neurological presentations described as dizziness, numbness, and forgetfulness, with the broad differential diagnoses that such symptoms encompass. Such self-induced symptoms may occur in the context of meditation and self-suggestion.

Reference

Cross References
Depersonalization; Dissociation

Derealization
Derealization, a form of dissociation, is the experience of feeling that the world around is unreal. There may be concurrent depersonalization.

Reference

Cross References
Depersonalization; Dissociation

Developmental Signs
- see FRONTAL RELEASE SIGNS; PRIMITIVE REFLEXES

Diagonistic Dyspraxia
A dissociative phenomenon observed after callosotomy, probably identical to intermanual conflict.

Reference

Cross References
Alien hand, Alien limb; Intermanual conflict

Diamond on Quadriceps Sign
Diamond on quadriceps sign may be seen in patients with dysferlinopathies (limb girdle muscular dystrophy type 2B, Miyoshi myopathy): with the knees slightly bent so that the quadriceps are in moderate action, an asymmetric diamond-shaped bulge may be seen, with wasting above and below, indicative of the selectivity of the dystrophic process in these conditions.
Diaphoresis

Diaphoresis is sweating, either physiological as in sympathetic activation (e.g. during hypotension, hypoglycaemia), or pathological (hyperhidrosis, q.v.). Diaphoresis may be seen in syncope, delirium tremens, or may be induced by certain drugs (e.g. cholinesterase inhibitors) or drug withdrawal (e.g. opiates in dependent individuals). Anticholinergics decrease diaphoresis but increase core temperature, resulting in a warm dry patient.

Cross Reference
Hyperhidrosis

Diaphragm Weakness

Diaphragm weakness is a feature of certain myopathies, such as acid maltase deficiency, and of cervical cord lesions (C3–C5) affecting phrenic nerve function. Forced vital capacity measured in the supine and sitting positions is often used to assess diaphragmatic function, a drop of 25% being taken as indicating diaphragmatic weakness.

Reference

Cross Reference
Paradoxical breathing

Diplophonia

Diplophonia, the simultaneous production of two pitch levels when phonating, occurs in unilateral vocal cord paralysis because each vocal fold has a different vibration frequency.

Cross References
Bovine cough; Dysphonia

Diplopia

Diplopia is double vision, viz., seeing two images of a single object. The spatial and temporal characteristics of the diplopia may help to ascertain its cause.

Diplopia may be monocular, in which case ocular causes are most likely (although monocular diplopia may be cortical or functional in origin), or binocular, implying a divergence of the visual axes of the two eyes. With binocular diplopia, it is of great importance to ask the patient whether the images are separated horizontally, vertically, or obliquely (tilted), since this may indicate the extraocular muscle(s) most likely to be affected. Whether the two images are
Diplopia

Diseparation or overlapping is important when trying to ascertain the direction of maximum diplopia.

The experience of diplopia may be confined to, or particularly noticeable during, the performance of particular activities, reflecting the effect of gaze direction; for example, diplopia experienced on coming downstairs may reflect a trochlear (IV) nerve palsy; or only on looking to the left may reflect a left abducens (VI) nerve palsy. Double vision experienced on looking at a distant object after looking down (e.g. reading) may occur with bilateral abducens (VI) nerve palsies. The effect of gaze direction on diplopia should always be sought, since images are most separated when looking in the direction of a paretic muscle. Conversely, diplopia resulting from the breakdown of a latent tendency for the visual axes to deviate (latent strabismus, squint) results in diplopia in all directions of gaze.

Examination of the eye movements should include asking the patient to look at a target, such as a pen, in the various directions of gaze (versions) to ascertain where diplopia is maximum. Ductions are tested monocularly with the opposite eye covered. Then, each eye may be alternately covered to try to demonstrate which of the two images is the false one, namely that from the non-fixing eye. The false image is also the most peripheral image. Thus in a left abducens (VI) nerve palsy, diplopia is maximum on left lateral gaze; when the normal right eye is covered the inner image disappears; the non-fixing left eye is responsible for the remaining false image, which is the more peripheral and which disappears when the left eye is covered.

Other clues to the cause of diplopia include ptosis (unilateral: oculomotor (III) nerve palsy; bilateral: myasthenia gravis), and head tilt or turn (e.g. turn to the right suggests a weak right lateral rectus muscle suggesting a right abducens (VI) nerve palsy; tilt to the left shoulder suggests a right trochlear (IV) nerve palsy, = Bielschowsky’s sign).

Manifest squints (heterotropia) are obvious but seldom a cause of diplopia if long-standing. Latent squints may be detected using the cover–uncover test, when the shift in fixation of the eyes indicates an imbalance in the visual axes; this may account for diplopia if the normal compensation breaks down. This produces diplopia in all directions of gaze (comitant). Patients may with an effort be able to fuse the two images.

Transient diplopia (minutes to hours) suggests the possibility of myasthenia gravis. There are many causes of persistent diplopia, including the breakdown of a latent strabismus, development of oculomotor (III), trochlear (IV), or abducens (VI) nerve palsy (singly or in combination), orbital myopathy (thyroid), and mass lesions of the orbit (tumour, pseudotumour).

Divergence of the visual axes or ophthalmoplegia without diplopia suggests a long-standing problem, such as amblyopia or chronic progressive external ophthalmoplegia. Some eye movement disorders are striking for the lack of associated diplopia, e.g. internuclear ophthalmoplegia.

References
Directional Hypokinesia

Directional hypokinesia is a reluctance to move towards contralesional space seen in the neglect syndrome.

Cross References
Motor neglect; Neglect

Disc Swelling
Swelling or oedema of the optic nerve head may be visualized by ophthalmoscopy. It produces haziness of the nerve fibre layer obscuring the underlying vessels; there may also be haemorrhages and loss of spontaneous retinal venous pulsation. Disc swelling due to oedema must be distinguished from pseudopapilloedema, elevation of the optic disc not due to oedema, in which the nerve fibre layer is clearly seen.

Disc swelling may be due to raised intracranial pressure (papilloedema, q.v.), or local inflammation of the optic nerve (papillitis), and may be associated with marked impairment of vision, for example, in optic neuritis, or be without specific visual complaint (as may be the case in papilloedema). The clinical history, visual acuity, and visual fields may help determine the cause of disc swelling. Recognized causes of disc swelling include

- **Unilateral:**
  - Optic neuritis
  - Acute ischaemic optic neuropathy (arteritic, non-arteritic)
  - Orbital compressive lesions, e.g. optic nerve sheath meningioma (Foster Kennedy syndrome)
  - Grave’s ophthalmopathy (through compression of retinal veins by myositis)
  - Central retinal vein occlusion
  - Infiltration: carcinoma, lymphoma, granuloma
  - Raised intracranial pressure (papilloedema; more usually bilateral)

- **Bilateral:**
  - Raised intracranial pressure (papilloedema)
  - Malignant hypertension
  - Hypercapnia
  - High CSF protein, as in Guillain–Barré syndrome
  - Any of the unilateral causes

Cross References
Foster Kennedy syndrome; Papilloedema; Pseudopapilloedema; Retinal venous pulsation; Visual field defects
Disinhibition
Disinhibited behaviour is impulsive, showing poor judgment and insight, and may transgress normal cultural or social bounds. There is a loss of normal emotional and/or behavioural control. The disinhibited patient may be inappropriately jocular (*witzelsucht*), short-tempered (verbally abusive, physically aggressive), distractible (impaired attentional mechanisms), and show emotional lability. A Disinhibition Scale encompassing various domains (motor, intellectual, instinctive, affective, sensitive) has been described.

Disinhibition is a feature of frontal lobe, particularly orbitofrontal, dysfunction. This may be due to neurodegenerative disorders (frontotemporal dementia, Alzheimer’s disease), mass lesions, or be a feature of epileptic seizures.

Cross References
Attention; Emotionalism, Emotional lability; Frontal lobe syndromes; *Witzelsucht*

Dissociated Sensory Loss
Dissociated sensory loss refers to impairment of selected sensory modalities with preservation, or sparing, of others. It is usually an indication of an intramedullary spinal cord lesion. For example, a focal central cord pathology such as syringomyelia will, in the early stages, selectively involve decussating fibres of the spinothalamic pathway within the ventral commissure, thus impairing pain and temperature sensation (often in a suspended, ‘cape-like’, ‘bathing suit’, ‘vest-like’, or cuirasse distribution), whilst the dorsal columns are spared, leaving proprioception intact. The anterior spinal artery syndrome also leaves the dorsal columns intact. Conversely, pathologies confined, largely or exclusively, to the dorsal columns (classically tabes dorsalis and subacute combined degeneration of the cord from vitamin B₁₂ deficiency, but probably most commonly seen with compressive cervical myelopathy) impair proprioception, sometimes sufficient to produce pseudoathetosis or sensory ataxia, whilst pain and temperature sensation is preserved. A double dissociation of sensory modalities on opposite sides of the trunk is seen in the Brown–Séquard syndrome.

Small fibre peripheral neuropathies may selectively affect the fibres which transmit pain and temperature sensation, leading to a glove-and-stocking impairment to these modalities. Neuropathic (Charcot) joints and skin ulceration may occur in this situation; tendon reflexes may be preserved.

Cross References
Analgesia; Ataxia; Brown–Séquard syndrome; Charcot joint; *Main succulente*; Myelopathy; Proprioception; Pseudoathetosis; Sacral sparing

Dissociation
Dissociation is an umbrella term for a wide range of symptoms involving feelings of disconnection from the body (depersonalization) or the surroundings (derealization). Common in psychiatric disorders (depression, anxiety, schizophrenia), these symptoms are also encountered in neurological conditions (epilepsy, migraine, presyncope), conditions such as functional weakness and non-epileptic attacks, and in isolation by a significant proportion of the general population. Symptoms of dizziness and blankness may well be the result of dissociative states rather than neurological disease.
Divisional Palsy

The oculomotor (III) nerve divides into superior and inferior divisions, usually at the superior orbital fissure. The superior division or ramus supplies the superior rectus and levator palpebrae superioris muscles; the inferior division or ramus supplies medial rectus, inferior rectus and inferior oblique muscles. Isolated dysfunction of these muscular groups allows diagnosis of a divisional palsy and suggests pathology at the superior orbital fissure or anterior cavernous sinus. However, occasionally this division may occur more proximally, at the fascicular level (i.e. within the midbrain) or within the subarachnoid space, giving a false-localizing divisional palsy. This may reflect the topographic arrangement of axons within the oculomotor nerve.

Reference
Larner AJ. Proximal superior division oculomotor nerve palsy from metastatic subarachnoid infiltration *Journal of Neurology* 2002; 249: 343–344.

Cross References
'False-localizing signs’; Oculomotor (III) nerve palsy

Dix–Hallpike Positioning Test
- see HALLPIKE MANOEUVRE, HALLPIKE TEST

Doll’s Eye Manoeuvre, Doll’s Head Manoeuvre

This test of the vestibulo-ocular reflex (VOR) is demonstrated by rotating the patient’s head and looking for a conjugate eye movement in the opposite direction. Although this can be done in a conscious patient focusing on a visual target, smooth pursuit eye movements may compensate for head turning; hence the head impulse test (q.v.) may be required. The manoeuvre is easier to do in the unconscious patient, when testing for the integrity of brainstem reflexes.

A slow (0.5–1.0 Hz) doll’s head manoeuvre may be used in conscious patients to assess vestibulo-ocular reflexes. Whilst directly observing the eyes, ‘catch up’ saccades may be seen in the absence of VOR. Measuring visual acuity with head movement compared to visual acuity with the head still (dynamic visual acuity, or illegible E test), two to three lines may be dropped if VOR is impaired. On ophthalmoscopy, the disc moves with the head if VOR is lost.

Reference
Roberts TA, Jenkyn LR, Reeves AG. On the notion of doll’s eyes. *Archives of Neurology* 1984; 41: 1242–1243.

Cross References
Bell’s phenomenon, Bell’s sign; Caloric testing; Coma; Head impulse test; Oculocephalic response; Supranuclear gaze palsy; Vestibulo-ocular reflexes
Dropped Head Syndrome

Dorsal Guttering
Dorsal guttering refers to the marked prominence of the extensor tendons on the dorsal surface of the hand when intrinsic hand muscles (especially interossei) are wasted, as may occur in an ulnar nerve lesion, a lower brachial plexus lesion, or a T1 root lesion. Benign extramedullary tumours at the foramen magnum may also produce this picture (remote atrophy, a ‘false-localizing sign’). In many elderly people the extensor tendons are prominent in the absence of significant muscle wasting.

Cross Reference
Wasting

Double Elevator Palsy
This name has been given to monocular elevation paresis. It may occur in association with pretectal supranuclear lesions either contralateral or ipsilateral to the paretic eye interrupting efferents from the rostral interstitial nucleus of the medial longitudinal fasciculus to the superior rectus and inferior oblique subnuclei. Bell’s phenomenon may be preserved.

Reference

Cross Reference
Bell’s phenomenon, Bell’s sign

Downbeat Nystagmus
- see NYSTAGMUS

Dressing Apraxia
- see APRAXIA

Drooling
- see SIALORRHOEA

Dropped Head Syndrome
Dropped head syndrome (head droop or head drop) refers to forward flexion of the head on the neck, such that the chin falls on to the chest (cf. antecollis) and the head cannot be voluntarily extended. This syndrome has a broad differential diagnosis, encompassing disorders which may cause axial truncal muscle weakness, especially of upper thoracic and paraspinal muscles.

- **Neuropathy/Neuronopathy:**
  - Motor neurone disease (the author has also seen this syndrome in a patient with frontotemporal lobar degeneration with motor neurone disease, FTLD/MND)
  - Guillain–Barré syndrome, chronic inflammatory demyelinating polyneuropathy
  - Paraneoplastic motor neuronopathy

- **Neuromuscular junction disorder:**
  - Myasthenia gravis
• **Myopathy:**
  - Polymyositis
  - Myotonic dystrophy
  - Myopathy with rimmed vacuoles
  - ‘Dropped head syndrome’, or ‘isolated neck extensor myopathy’, a condition of uncertain aetiology but which may on occasion be steroid-responsive (‘bent spine syndrome’ or camptocormia may be a related form of axial myopathy)

• **Extrapyramidal disorders:**
  - Parkinson’s disease
  - Multiple system atrophy
  - Progressive supranuclear palsy

  Of these, probably MND and myasthenia gravis are the most common causes.

  Treatment of the underlying condition may be possible, hence investigation is mandatory. If not treatable (e.g. MND), a head brace may keep the head upright.

**References**


**Cross References**

Antecollis; Camptocormia; Myopathy

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**Drusen**

Drusen are hyaline bodies that are typically seen on and around the optic nerve head and may be mistaken for papilloedema (‘pseudopapilloedema’). Drusen are thought to result from altered axonal flow with axonal degeneration. They occur sporadically or may be inherited in an autosomal dominant fashion, and are common, occurring in 2% of the population. In children the drusen are buried whilst in adults they are on the surface.

Drusen are usually asymptomatic but can cause visual field defects (typically an inferior nasal visual field loss) or occasionally transient visual obscurations, but not changes in visual acuity; these require investigation for an alternative cause. When there is doubt whether papilloedema or drusen is the cause of a swollen optic nerve head, retinal fluorescein angiography is required.

**Reference**


**Cross References**

Disc swelling; Papilloedema; Pseudopapilloedema; Visual field defects
**Dynamic Aphasia**

Dynamic aphasia refers to an aphasia characterized by difficulty initiating speech output, ascribed to executive dysfunction. There is a reduction in spontaneous speech, but on formal testing there are no paraphasias, minimal anomia, preserved repetition, and automatic speech. ‘Incorporational echolalia’, when the patient uses the examiner’s question to help form an answer, may be observed. Dynamic aphasia may be conceptualized as a variant of transcortical motor aphasia and may be seen with lesions of dorsolateral prefrontal cortex (‘frontal aphasia’). It has also been reported in progressive supranuclear palsy. A division into pure and mixed forms has been suggested, with additional phonological, lexical, syntactical, and articulatory impairments in the latter.

**Reference**


**Cross References**

Echolalia; Transcortical aphasias

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**Dysaesthesia**

Dysaesthesia is an unpleasant, abnormal or unfamiliar, sensation, often with a burning and/or ‘electrical’ quality. Some authorities reserve the term for provoked positive sensory phenomena, as opposed to spontaneous sensations (paraesthesia). Dysaesthesia differs from paraesthesia in its unpleasant quality, but may overlap in some respects with allodynia, hyperalgesia, and hyperpathia (the latter phenomena are provoked by stimuli, either non-noxious or noxious).

There are many causes of dysaesthesia, both peripheral (including small fibre neuropathies, neuroma, and nerve trauma) and central (e.g. spinal multiple sclerosis).

Dysaesthetic sensations may be helped by agents such as carbamazepine, amitriptyline, gabapentin, and pregabalin.

**Cross References**

Allodynia; Hyperalgesia; Hyperpathia; Paraesthesia

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**Dysarthria**

Dysarthria is a disorder of speech, as opposed to language (cf. aphasia), because of impairments in the actions of the speech production apparatus per se, due to paralysis, ataxia, tremor, or spasticity, in the presence of intact mental function, comprehension, and memory of words. In its most extreme form, anarthria, there is no speech output.

Dysarthria is a symptom, which may be caused by a number of different conditions, all of which ultimately affect the function of pharynx, palate, tongue, lips, and larynx, be that at the level of the cortex, lower cranial nerve nuclei or their motor neurones, neuromuscular junction, or bulbar muscles themselves. Dysarthrias affect articulation in a highly reliable and consistent manner, the errors reflecting the muscle group involved in the production of specific sounds. There are various syndromes of dysarthria, which have been classified as follows:
Flaccid or nasal dysarthria:
hypernasal, breathy, whining output, as in bulbar palsy, e.g. myasthenia gravis.

Spastic dysarthria:
slow, strained (‘strangled’) output, monotonous, as in pseudobulbar palsy; may coexist with Broca’s aphasia.

Ataxic or cerebellar dysarthria:
altered rhythm of speech, uneven irregular output, slurred speech (as if inebriated), improper stresses; seen in acute cerebellar damage due to asynergia of speech muscle contractions (cf. scanning speech).

Hypokinetic dysarthria:
monotonic pitch, hypophonic volume, as in parkinsonism.

Hyperkinetic dysarthria:
several varieties are described, including choreiform (as in Huntington’s disease), dystonic (as in tardive dyskinesia, and other dystonic syndromes), tremulous (tremor syndromes), and the dysarthria with vocal tics (including coprolalia) in Tourette syndrome.

Mixed dysarthria:
combination of any of above.

Recognized causes of dysarthria include the following:

- Muscle disease:
e.g. oculopharyngeal muscular dystrophy: nasal speech; weak pharynx/drooling.

- Neuromuscular disorder:
e.g. myasthenia gravis: nasal speech; fatiguability (development of hypophonia with prolonged conversation, counting).

- Lower motor neurone disease = bulbar palsy:
e.g. motor neurone disease (rasping monotones, wasted and fasciculating tongue), poliomyelitis, Guillain–Barré syndrome, diphtheria.

- Upper motor neurone disease = pseudobulbar palsy:
e.g. motor neurone disease (spastic tongue), cerebrovascular disease.

- Cortical dysarthria:
damage to left frontal cortex, usually with associated right hemipareses; may be additional aphasia.

- Extrapyramidal disease:
e.g. hypokinetic disorders: Parkinson’s disease: slow, hypophonic, monotonic; multiple system atrophy (may have vocal cord palsy).
e.g. hyperkinetic disorders: Huntington’s disease: loud, harsh, variably stressed, and poorly coordinated with breathing; myoclonus of any cause (hiccup speech); dystonia of any cause.

- Ataxic dysarthria:
disease of or damage to cerebellum: slow, slurred, monotonous, with incoordination of speech with respiration; may therefore be quiet
Dysexecutive Syndrome

and then explosive; unnatural separation of syllables; slow tongue movements.

- Acquired stuttering:
  involuntary repetition of letters or syllables, may be acquired with aphasia; developmental stutter, the more common cause, usually affects the beginnings of words and with plosive sounds, whereas the acquired form may be evident throughout sentences and affect all speech sounds.

Treatment of the underlying cause may improve dysarthria (e.g. nasal dysarthria of myasthenia gravis). Baclofen has been suggested for dysarthria of upper motor neurone type. Speech and language therapy may provide symptomatic benefit.

References

Cross References
Anarthria; Aphasia; Asynergia; Broca’s aphasia; Bulbar palsy; Coprolalia; Dysphonia; Fatigue; Lower motor neurone (LMN) syndrome; Parkinsonism; Pseudobulbar palsy; Scanning speech; Stutter; Upper motor neurone (UMN) syndrome

Dyscalculia
- see ACALCULIA

Dyschromatopsia
- see ACHROMATOPSIA

Dysdiadochokinesia
Dysdiadochokinesia or adiadochokinesia is a difficulty in performing rapid alternating movements, for example, pronation/supination of the arms, tapping alternately with the palm and dorsum of the hand, tapping the foot on the floor.

Dysdiadochokinesia is a sign of cerebellar dysfunction, especially hemisphere disease, and may be seen in association with asynergia, ataxia, dysmetria, and excessive rebound phenomenon. It may reflect the impaired checking response seen in cerebellar disease. Dysdiadochokinesia may also be seen with disease of the frontal lobes (‘frontal apraxia’) or basal ganglia.

Cross References
Asynergia; Apraxia; Ataxia; Cerebellar syndromes; Dysmetria; Rebound phenomenon

Dysexecutive Syndrome
The term executive function encompasses a range of cognitive processes including sustained attention, fluency and flexibility of thought, problem-solving skills,
and planning and regulation of adaptive and goal-directed behaviour. Some authors prefer to use these individual terms, rather than ‘lump’ them together as executive function. Deficits in these various functions, the dysexecutive syndrome, are typically seen with lateral prefrontal cortex lesions.

**Reference**

**Cross References**
Attention; Frontal lobe syndromes

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**Dysgeusia**
Dysgeusia is a complaint of distorted taste perception. It may occur along with anosmia as a feature of upper respiratory tract infections and has also been described with various drug therapies, in psychiatric diseases, and as a feature of zinc deficiency.

**Reference**

**Cross References**
Ageusia; Anosmia

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**Dysgraphaesthesia**
- see AGRAPHOGNOSIA; GRAPHAESTHESIA

**Dysgraphia**
- see AGRAPHIA

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**Dyskinesia**
Dyskinesia may be used as a general term for excessive involuntary movements, encompassing tremor, myoclonus, chorea, athetosis, tics, stereotypies, and hyperkplexia. The term may be qualified to describe a number of other syndromes of excessive movement, e.g.:

- **Drug-induced dyskinesia:**
  Fluid, restless, fidgety movements seen in patients with Parkinson’s disease after several years of levodopa therapy, and often described according to their relationship to timing of tablets (e.g. peak dose, diphasic), although others are unpredictable (freezing, yo-yo-ing). In MPTP-induced parkinsonism, dyskinesias tend to occur early, hence it may be the depth of dopamine deficiency rather than chronicity of treatment which is the key determinant; reduction in overall levodopa use (increased frequency of smaller doses, controlled-release preparations, addition of dopamine agonists) may reduce these effects; amantadine is sometimes helpful.

- **Tardive dyskinesia:**
  A form of drug-induced dyskinesia developing after long-term use of neuroleptic (dopamine antagonist) medication, typically involving...
Dysmetria

Dysmetria, or past-pointing, is a disturbance in the control of range of movement in voluntary muscular action and is one feature of the impaired checking response seen in cerebellar lesions (especially cerebellar hemisphere lesions).

Dysmetria may also be evident in saccadic eye movements: hypometria (undershoot) is common in parkinsonism; hypermetria (overshoot) is more typical of cerebellar disease (lesions of dorsal vermis and fastigial nuclei).

In cerebellar disorders, dysmetria reflects the asynergia of coordinated muscular contraction.

References

Dyslexia

Dyslexia is difficulty or impairment in reading, usually applied to developmental abnormalities of reading ability. A loss of previously acquired reading ability is probably better termed alexia.

Cross Reference
Alexia

Dysmentia

The term dysmentia has been suggested as an alternative to dementia, to emphasize the possibility of treating and preventing cognitive decline.

Reference

Cross Reference
Dementia

Dysmetria

orolinguial musculature (buccolinguual syndrome, rabbit syndrome, ‘bon-bon sign’) and occasionally trunk and arms; usually persists after withdrawal of causative therapy; clonazepam, baclofen, and tetrabenazine may help.

- Paroxysmal dyskinesias:
  Paroxysmal kinesigenic choreoathetosis/dystonia (PKC; usually responds to carbamazepine) and paroxysmal non-kinesigenic dystonia/choreoathetosis (PDC; does not respond to carbamazepine).

- Focal dyskinesias:
  Oro-facial dyskinesia, belly dancer’s dyskinesia, moving ear syndrome.

References

Cross References
Athetosis; ‘Bon-bon sign’; Chorea, Choreoathetosis; Dystonia; Hyperekplexia; Moving ear; Myclonus; Parkinsonism; Stereotypy; Tic; Yo-yo-ing

Dyslexia

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Cross Reference
Alexia

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Dementia

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In cerebellar disorders, dysmetria reflects the asynergia of coordinated muscular contraction.

References

**Cross References**
Asynergia; Cerebellar syndromes; Dysdiadochokinesia; Parkinsonism; Rebound phenomenon; Saccades

**Dysmorphopsia**
The term dysmorphopsia has been proposed for impaired vision for shapes, a visual recognition defect in which visual acuity, colour vision, tactile recognition, and visually guided reaching movements are intact. These phenomena have been associated with bilateral lateral occipital cortical damage (e.g. after carbon monoxide poisoning) and are thought to reflect a selective loss of the magnocellular visual pathway. Whether this condition is an agnosia for shape or visual form, or a perceptual problem (‘pseudoagnosia’), remains a subject of debate and the term dysmorphopsia has been suggested as a compromise between the different strands of thought.

**Reference**

**Cross References**
Agnosia; Visual agnosia; Visual form agnosia

**Dysnomia**
- see ANOMIA

**Dysphagia**
Dysphagia is difficulty swallowing. This may have local mechanical causes which are usually gastroenterological in origin (tumour; peptic ulceration/stricture, in which case there may be additional pain on swallowing – odynophagia) but sometimes vascular (aberrant right subclavian artery – dysphagia lusoria) or due to connective tissue disease (systemic sclerosis). Dysphagia of neurological origin may be due to pathology occurring anywhere from cerebral cortex to muscle. Neurological control of swallowing is bilaterally represented and so unilateral upper motor neurone lesions may cause only transient problems. Poststroke dysphagia is common, but there is evidence of cortical reorganization (neuroplasticity) underpinning recovery. Bilateral upper motor neurone lesions cause persistent difficulties. Dysphagia of neurological origin may be accompanied by dysphonia, palatal droop, and depressed or exaggerated gag reflex.

Dysphagia may be
- **Neurogenic**:
  - CNS:
    - Cerebrovascular disease: hemisphere, brainstem stroke
    - Extrapyramidal disease: Parkinson’s disease, progressive supranuclear palsy, Huntington’s disease, Wilson’s disease, tardive dyskinesia, dystonia
    - Inflammatory disease: multiple sclerosis
Dysphagia

Neoplasia: primary, secondary; cerebral, brainstem (skull base)
Other structural disorders of the brainstem: syringobulbia, cerebellar disease
Developmental disorders: cerebral palsy syndromes, Chiari malformations
Neuronopathy:
   Motor neurone disease
Neuropathy:
   Guillain–Barré syndrome
   Autonomic neuropathy (diabetes mellitus, amyloidosis, Chagas’ disease, autonomic failure, Riley Day syndrome)
   Lower motor neurone pathology: bulbar palsy, isolated vagus (X) nerve palsy, jugular foramen syndrome
Neuromuscular:
   Myasthenia gravis
   • Myogenic:
      Inflammatory muscle disease: polymyositis, inclusion body myositis
      Myotonia: myotonic dystrophy
      Muscular dystrophy: oculopharyngeal muscular dystrophy
      Symptomatic oesophageal peristalsis (‘nutcracker oesophagus’)
   • Functional:
      ‘Hysterical’, globus hystericus (diagnosis of exclusion)
Gastrointestinal causes of dysphagia include
   • Intrinsic:
      Oesophageal carcinoma
      Metastatic or extrinsic tumour spread
      Peptic (postinflammatory) stricture
      Hiatus hernia
   • Extrinsic:
      Thoracic aortic aneurysm
      Abnormal origin of right subclavian artery (dysphagia lusoria)
      Posterior mediastinal mass
      Large goitre
      Retropharyngeal mass

If swallowing is compromised with a risk of aspiration, feeding may need to be undertaken via nasogastric tube, percutaneous gastrostomy or jejunostomy placed endoscopically (PEG or PEJ), or even parenterally.

References

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Dysphasia

Cross References
Bulbar palsy; Dysphonia; Gag reflex; Jugular foramen syndrome; Pseudobulbar palsy

Dysphasia
'The inaccuracy of applying an absolute negation [i.e. aphasia] to a partial defect [of language] has led to the suggestion of ‘dysphasia’ as a frequent substitute. The term does not, however, seem likely to come into use, a matter of little regret, since the word has not the merit of unimpeachable exactness, and it has an unfortunate resemblance in sound to ‘dysphagia’.'

Reference

Dysphonia
Dysphonia is a disorder of the volume, pitch, or quality of the voice resulting from dysfunction of the larynx, i.e. a disorder of phonation or sound generation. Hence this is a motor speech disorder and could be considered as a dysarthria if of neurological origin.

Dysphonia manifests as hoarseness or a whispering breathy quality to the voice. Diplophonia may occur. At the extreme, there may be complete loss of the voice (aphonia).

Recognized causes of dysphonia include
- Infection (laryngitis);
- Structural abnormalities, e.g. polyp, nodule, papilloma of vocal cord;
- Neurological causes:
  - Focal dystonic syndrome: spasmodic dysphonia or laryngeal dystonia (either abductor or adductor); the voice may have a strained and harsh quality, with low volume and pitch, vocal tremor, and irregularly distributed stoppages; with continuing speech, or if holding a single note, the voice may fade away entirely. These syndromes may be amenable to treatment with botulinum toxin.
  - Flaccid dysphonia, due to superior laryngeal nerve or vagus nerve (recurrent laryngeal nerve) palsy, bulbar palsy.

Reference

Dyspraxia
Dyspraxia is difficulty or impairment in the performance of a voluntary motor act despite an intact motor system and level of consciousness. This may be developmental in origin (‘clumsy child’), but in adult practice reflects a loss of function (hence apraxia is a better term).
**Dystonia**

Dystonia, a term first used by Oppenheim in 1911, is a motor syndrome of sustained involuntary muscle contractions causing twisting and repetitive movements, sometimes tremor, and/or abnormal postures. Dystonic movements may initially appear with voluntary movement of the affected part (‘action dystonia’) but may eventually occur with voluntary movement elsewhere in the body (‘overflow’). The severity of dystonia may be reduced by sensory tricks (*geste antagoniste*), using tactile or proprioceptive stimuli to lessen or eliminate posturing; this feature is unique to dystonia. Dystonia may develop after muscle fatiguing activity, and patients with focal dystonias show more rapid fatigue than normals. Dystonic disorders may be classified according to:

- **Age of onset**: the most significant predictor of prognosis: worse with earlier onset.
- **Distribution**: focal, segmental, multifocal, generalized; hemidystonia.
- **Aetiology**: primary/idiopathic vs. secondary/symptomatic.

Primary/idiopathic dystonias include the following:

- Primary torsion dystonia (idiopathic torsion dystonia);
- Severe generalized dystonia (dystonia musculorum deformans);
- Segemental, multifocal, and focal dystonias (e.g. torticollis, blepharospasm, writer’s cramp);
- Dopa-responsive dystonia (DRD; Segawa’s syndrome);
- Myoclonic dystonia.

Secondary/symptomatic dystonia: the differential diagnosis is broad, with more than 40 known causes, including

- Heredodegenerative disorders: Wilson’s disease, Huntington’s disease, neurodegeneration with brain iron accumulation, mitochondrial disorders, X-linked dystonia–parkinsonism (lubag);
- Paroxysmal dystonias/dyskinesias: paroxysmal kinesigenic choreoathetosis/dystonia (PKC; usually responds to carbamazepine) and paroxysmal non-kinesigenic dystonia/choreoathetosis (PDC; does not respond to carbamazepine);
- Metachromatic leucodystrophy;
- Gangliosidoses (GM1, GM2);
- Perinatal cerebral injury;
Dystonia

• Encephalitis;
• Head trauma;
• Multiple sclerosis;
• Drugs/toxins, e.g. antipsychotic, antiemetic, and antidepressant drugs;
• Psychogenic.

Appropriate investigations to exclude these symptomatic causes (especially Wilson’s disease) are appropriate.

The pathogenesis of dystonia is incompletely understood. Different mechanisms may apply in different conditions. Peripheral focal dystonias such as torticollis and writer’s cramp have been suggested to result from abnormal afferent information relayed from ‘stiff’ muscle spindles. The genetic characterization of various dystonic syndromes may facilitate understanding of pathogenesis.

From the therapeutic point of view, one of the key questions relates to response to levodopa: dopa-responsive dystonia (DRD) responds very well to levodopa (and response fluctuations do not develop over time; cf. Parkinson’s disease). Other treatments which are sometimes helpful include anticholinergics, dopamine antagonists, dopamine agonists, and baclofen. Drug-induced dystonia following antipsychotic, antiemetic, or antidepressant drugs is often relieved within 20 min by intramuscular biperiden (5 mg) or procyclidine (5 mg). Botulinum toxin may be very helpful in some focal dystonias (e.g. blepharospasm). Surgery for dystonia using deep brain stimulation is still at the experimental stage.

References

Cross References
Anismus; Blepharospasm; Dysphonia; Eyelid apraxia; Fatigue; Gaping; Geste antagoniste; Hemidystonia; Torticollis; Writer’s cramp
Ear Click
- see PALATAL MYOCLONUS; TINNITUS

Eastchester Clapping Sign
The Eastchester clapping sign is advocated as an early and sensitive sign of hemispatial egocentric neglect. Patients are asked to clap: those with neglect perform one-handed motions which stop at the midline. Hemiplegic patients without neglect reach across the midline and clap against their plegic hand.

Reference

Cross-Reference
Neglect

Echolalia
Echolalia is the involuntary automatic repetition of an interviewer’s speech. This may be observed in a variety of clinical situations:

- *Transcortical sensory aphasia:*
  In the context of a fluent aphasia with repetition often well or normally preserved, usually as a result of a vascular lesion of the left hemisphere although an analogous situation may be encountered in Alzheimer’s disease; ‘incorporational echolalia’, when the patient uses the examiner’s question to help form an answer, may be observed as a feature of ‘dynamic aphasia’ which bears resemblance to transcortical motor aphasia, but may result from a frontal lesion.

- *Transcortical motor aphasia:*
  ‘Effortful echolalia’ has been reported in the context of infarction of the left medial frontal lobe, including the supplementary motor area, showing that neither the ability to repeat nor fluent speech is required for echolalia.

- *Tourette syndrome:*
  As a complex vocal tic, along with coprolalia.

- *Alzheimer’s disease, frontotemporal lobar degeneration:*
  As a symptom of cognitive impairment/dementia.

- *Epilepsy:*
  From a left frontal lobe focus.

- *Schizophrenia:*
  As a catatonic symptom.
• *Early infantile autism, mental retardation:*  
  As a reflection of pathological mental development.

• *Frontal lobe lesions:*  
  As a feature of imitation behaviour.

• *Normal children:*  
  At a particular stage of language acquisition.

**References**


**Cross References**

Aphasia; Coprolalia; Dynamic aphasia; Imitation behaviour; Jargon aphasia; Logorrhoea; Palilalia; Transcortical aphasias

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**Echophenomena**

- see IMITATION BEHAVIOUR

**Echopraxia**

Echopraxia is the involuntary, automatic, imitation of an interviewer’s movements. This may be observed as a feature of apraxic syndromes such as corticobasal degeneration, as a complex motor tic in Tourette syndrome, and in frontal lobe disorders (imitation behaviour).

**Reference**


**Cross References**

Copropraxia; Imitation behaviour; Tic

**Écriture en Double Miroir**

- see MIRROR WRITING

**Ectropion**

- see LID RETRACTION

**Eidetic Memory**

Photographic, or eidetic, memory is an enhancement of memory to prodigious capacity, beyond hypermnesia. Synaesthesia may be linked to eidetic memory; synaesthesia being used as a mnemonic aid.

**Reference**


**Cross Reference**

Synaesthesia

**Eight-and-a-Half Syndrome**

The combination of a facial (VII) nerve palsy with a one-and-a-half syndrome due to a pontine lesion has been labelled the eight-and-a-half syndrome. Patients
may develop oculopalatal myoclonus months to years after the onset of the ocular motility problem.

**References**


**Cross References**

Facial paresis, Facial weakness; Myoclonus; One-and-a-half syndrome; Palatal myoclonus

**Ekbom’s Syndrome**

Patients with Ekbom’s syndrome or delusional parasitosis believe with absolute certainty that insects, maggots, lice, or other vermin infest their skin or other parts of the body. Sometimes other psychiatric features may be present, particularly if the delusions are part of a psychotic illness such as schizophrenia or depressive psychosis. Females are said to be more commonly affected. Clinical examination may sometimes show evidence of skin picking, scratching, or dermatitis caused by repeated use of antiseptics. The patient may produce skin fragments or other debris as ‘evidence’ of infestation. Treatment should be aimed at the underlying condition if appropriate; if the delusion is isolated, antipsychotics such as pimozide may be tried.

**Reference**


**Emotionalism, Emotional Lability**

Emotionalism, or emotional lability, or emotional incontinence, implies both frequent and unpredictable changes in emotional expression, for example, tearfulness followed shortly by elation, and an inappropriate expression of emotion, for example, uncontrollable (‘uninhibited’ or disinhibited) laughter or crying. A distinction may be drawn between the occurrence of these phenomena spontaneously or without motivation, or in situations which although funny or sad are not particularly so. Also, a distinction may be made between such phenomena when there is congruence of mood and affect, sometimes labelled with terms such as moria or *witzelsucht* (e.g. laughing when feeling happy or elated), and when there is no such congruence (e.g. laughing when not feeling happy or elated), sometimes labelled as pathological, forced, or inappropriate laughter and crying.

The neurobehavioural state of emotional lability reflects frontal lobe (especially orbitofrontal) lesions, often vascular in origin, and may coexist with disinhibited behaviour. It is more common in vascular dementia than in Alzheimer’s disease. It may also be seen in delirium and in psychiatric disorders (mania). Pathological laughter and crying may occur as one component of pseudobulbar palsy (‘pseudobulbar affect’).
Emposthotonos

Emposthotonos is an abnormal posture consisting of flexion of the head on the trunk and the trunk on the knees, sometimes with flexion of the limbs (cf. opisthotonos). Such attacks of ‘bowing’ may be seen in infantile epilepsy syndromes such as West’s syndrome, sometimes called salaam seizures or jack-knife spasms.

Cross References
Opisthotonos; Seizures; Spasm

Encephalopathy

Encephalopathy is a general term referring to any acute or chronic diffuse disturbance of brain function. Characteristically it is used to describe an altered level of consciousness, which may range from drowsiness to a failure of selective attention, to hypervigilance; with or without: disordered perception, memory (i.e. cognitive deficits); epileptic seizures; headache; abnormal movements such as tremor, myoclonus, or asterixis; and focal neurological deficits (less common). Clearly, these features overlap with those of delirium.

As with terms such as coma and stupor, it is probably better to give a description of the patient’s clinical state rather than use a term that is open to variable interpretation. Although the term encephalopathy is sometimes reserved for metabolic causes of diffuse brain dysfunction, this usage is not universal. Conditions which may be described as an encephalopathy include

- **Metabolic disorders**: hypoxia/ischaemia, hypoglycaemia; organ failure, electrolyte disturbances, hypertension;
- Drug/toxin ingestion;
- Brain inflammation/infection (e.g. encephalitis);
- Miscellaneous conditions, e.g. Alzheimer’s disease, Creutzfeldt–Jakob disease.

Cross References
Asterixis; Coma; Delirium; Myoclonus; Stupor; Tremor

En Garde Position

- see FENCER’S POSTURE, FENCING POSTURE

Enophthalmos

Enophthalmos is an inward displacement of the eyeball (sinking or withdrawal) into the eye socket (cf. exophthalmos). It is classically described as one of the cardinal features of Horner’s syndrome (along with miosis, ptosis, and anhidrosis) but is seldom actually measured. Enophthalmos may also occur in dehydration (probably the most common cause), orbital trauma (e.g. orbital...
floor fracture), senile orbital fat atrophy, hemifacial atrophy, and orbital tumour causing tethering and posterior traction on the eyeball.

**Cross References**
Anhidrosis; Exophthalmos; Hemifacial atrophy; Horner’s syndrome; Miosis; Ptosis

**Entomopia**
Entomopia (literally ‘insect eye’) is the name given to a grid-like pattern of multiple copies of the same visual image; hence, this is a type of polyopia. This phenomenon has been reported in migraine; its pathogenesis is uncertain.

**References**

**Cross Reference**
Polyopia

**Environmental Dependency Syndrome**
- see IMITATION BEHAVIOUR; UTILIZATION BEHAVIOUR

**Environmental Tilt**
Environmental tilt, also known as tortopia, is the sensation that visual space is tilted on its side or even upside down (‘floor-on-ceiling’ phenomenon, ‘upside-down’ reversal of vision, *verkehrtessehen*). This may last seconds to minutes. The temptation to dismiss such bizarre symptoms as functional should be resisted, since environmental tilt is presumed to reflect damage to connections between cerebellar and central vestibular-otolith pathways. It has been reported in the following situations:

- Lateral medullary syndrome of Wallenberg
- Transient ischaemic attacks in basilar artery territory
- Demyelinating disease
- Head injury
- Encephalitis
- Following third ventriculostomy for hydrocephalus

**Cross References**
Lateral medullary syndrome; Vertigo; Vestibulo-ocular reflexes

**Epiphora**
Epiphora is overflow of tears down the cheek. This may be not only due to a blocked nasolacrimal duct, or irritation to the cornea causing increased lacrimation, but it may also be neurological in origin, e.g. due to the sagging of the lower eyelid (ectropion) in a peripheral facial (VII) nerve (Bell’s) palsy, or the ‘crocodile tears’ following aberrant facial nerve regeneration. Lacrimation is also a feature of trigeminal autonomic cephalalgias such as cluster headache.

**Cross References**
Bell’s palsy; Crocodile tears
Epley Manoeuvre
- see HALLPIKE MANOEUVRE, HALLPIKE TEST; VERTIGO

Erythropsia
This name has been given to a temporary distortion of colour vision in which objects take on an abnormal reddish hue. This has been characterized as a visual illusion. There are various causes, including drug use, visual diseases, and pseudophakia.

Cross References
Illusion; ‘Monochromatopsia’; Phantom chromatopsia

Esophoria
Esophoria is a variety of heterophoria in which there is a tendency for the visual axes to deviate inward (latent convergent strabismus). Clinically this may be observed using the cover–uncover test as an outward movement of the covered eye as it is uncovered. Esophoria may occur in individuals with hyperopia (long-sightedness).

Cross References
Cover tests; Exophoria; Heterophoria

Esotropia
Esotropia is a variety of heterotropia in which there is manifest inward turning of the visual axis of one eye; the term is synonymous with convergent strabismus. It may be demonstrated using the cover test as an outward movement of the eye which is forced to assume fixation by occlusion of the other eye.

Esotropia may be associated with congenital latent nystagmus (i.e. nystagmus appearing when one eye is covered) in the presence of amblyopia; the slow phase in the viewing eye is towards the nose.

With lateral rectus muscle paralysis, the eyes are esotropic or crossed on attempted lateral gaze towards the paralyzed side, but the images are uncrossed. Acute esotropia has been described following contralateral thalamic infarction.

Cross References
Amblyopia; Cover tests; Diplopia; Exotropia; Heterotropia; Nystagmus

Eutonia
Kinnier Wilson used this term to describe an emotional lack of concern associated with the dementia of multiple sclerosis. It may perhaps reflect the cognitive anosognosia of a dementia syndrome.

Ewart Phenomenon
This is the elevation of ptotic eyelid on swallowing, a synkinetic movement. The mechanism is said to be aberrant regeneration of fibres from the facial (VII) nerve to the oculomotor (III) nerve innervating the levator palpebrae superioris muscle.

Cross References
Ptosis; Synkinesia, Synkinesis
**Exophoria**
Exophoria is a variety of heterophoria in which there is a tendency for the visual axes to deviate outward (latent divergent strabismus). Clinically this may be observed in the cover–uncover test as an inward movement as the covered eye is uncovered. Exophoria may occur in individuals with myopia and may be physiological in many subjects because of the alignment of the orbits.

**Cross References**
Cover tests; Esophoria; Heterophoria

**Exophthalmos**
Exophthalmos is forward displacement of the eyeball. The definition and the causes overlap with proptosis. The most common cause is dysthyroid eye disease (Graves’ disease).

**Cross References**
Lid retraction; Proptosis

**Exosomaesthesia**
The sensory disturbance associated with parietal lobe lesions may occasionally lead the patient to refer the source of a stimulus to some point outside the body, exosomaesthesia. A possible example occurs in Charles Dickens’s novel *Hard Times* (1854) in which Mrs Gradgrind locates her pain as ‘somewhere in the room’.

**Exotropia**
Exotropia is a variety of heterotropia in which there is manifest outward turning of the visual axis of an eye; the term is synonymous with divergent strabismus. It may be demonstrated using the cover test as an inward movement of the eye which is forced to assume fixation by occlusion of the other eye.

When the medial rectus muscle is paralyzed, the eyes are exotropic (wall-eyed) on attempted lateral gaze towards the paralyzed side, and the images are crossed.

**Cross References**
Cover tests; Esotropia; Heterotropia

**Extensor Posturing**
- see DECELERATE RIGIDITY

**External Malleolar Sign**
- see CHADDOCK’S SIGN

**External Ophthalmoplegia**
- see OPHTHALMOPARESIS, OPHTHALMOPLEGIA

**Extinction**
Extinction is the failure to respond to a novel or meaningful sensory stimulus on one side when a homologous stimulus is given simultaneously to the contralateral side (i.e. double simultaneous stimulation); it is sometimes called ‘suppression’. The stimuli may be visual, auditory, or tactile, e.g. asking the patient to say
which hand is touched when the eyes are shut. It is important to show that the patient responds appropriately to each hand being touched individually, but then neglects one side when both are touched simultaneously. With repeated testing the phenomenon may break down: extinguishing of extinction.

More subtle defects may be tested using simultaneous bilateral heterologous (asymmetrical) stimuli, although it has been shown that some normal individuals may show extinction in this situation.

A motor form of extinction has been postulated, manifesting as increased limb akinesia when the contralateral limb is used simultaneously.

The presence of extinction is one of the behavioural manifestations of neglect and most usually follows non-dominant (right) hemisphere (parietal lobe) lesions. There is evidence for physiological interhemispheric rivalry or competition in detecting stimuli from both hemifields, which may account for the emergence of extinction following brain injury.

Reference

Cross References
Akinesia; Hemiakinesia; Neglect; Visual extinction

**Extrapyramidal Signs**
- see PARKINSONISM

**Eyelid Apraxia**
Eyelid apraxia is an inability to open the eyelids at will, although they may open spontaneously at other times (i.e. voluntary–automatic dissociation). Eyelids may be opened manually or by a backwards head thrust. The term has been criticized on the grounds that this may not always be a true ‘apraxia’, in which case the term ‘levator inhibition’ may be preferred since the open eyelid position is normally maintained by tonic activity of the levator palpebrae superioris. Clinically there is no visible contraction of orbicularis oculi, which distinguishes eyelid apraxia from blepharospasm (however, perhaps paradoxically, the majority of cases of eyelid apraxia occur in association with blepharospasm). Neurophysiological studies do in fact show abnormal muscle contraction in the pretarsal portion of orbicularis oculi, which has prompted the suggestion that ‘focal eyelid dystonia’ may be a more appropriate term. Although the phenomenon may occur in isolation, associations have been reported with:
- Progressive supranuclear palsy (Steele–Richardson–Olszewski syndrome)
- Parkinson’s disease
- Huntington’s disease
- Multiple system atrophy
- MPTP intoxication
- Motor neurone disease
- Acute phase of non-dominant hemisphere cerebrovascular event
- Wilson’s disease
- Neuroacanthocytosis
The precise neuroanatomical substrate is unknown but the association with basal ganglia disorders points to involvement of this region. The underlying mechanisms may be heterogeneous, including involuntary inhibition of levator palpebrae superioris. Botulinum toxin injections may be helpful in some patients.

References

Cross References
Apraxia; Blepharospasm; Dystonia
Face–Hand Test
- see ‘ARM DROP’

Facial Paresis, Facial Weakness
Facial paresis, or prosopoplegia, may result from:

- Central (upper motor neurone) lesions
- Peripheral (lower motor neurone; facial (VII) nerve) lesions
- Neuromuscular junction transmission disorders
- Primary disease of muscle (i.e. myogenic)

Facial paresis is clinically heterogeneous which may be helpful with lesion localization.

- **Upper motor neurone facial weakness** (‘central facial palsy’):
  The ability to raise the eyebrow is preserved due to bilateral supranuclear connections to the frontalis muscle. A dissociation between volitional and emotional facial movements may also occur. Emotional facial palsy refers to the absence of emotional facial movement but with preserved volitional movements, as may be seen with frontal lobe (especially non-dominant hemisphere) precentral lesions (as in abulia, Fisher’s sign) and in medial temporal lobe epilepsy with contralateral mesial temporal sclerosis. Volitional paresis without emotional paresis may occur when corticobulbar fibres are interrupted (precentral gyrus, internal capsule, cerebral peduncle, upper pons).

Causes of upper motor neurone facial paresis include

**Unilateral:**
- Hemisphere infarct (with hemiparesis)
- Lacunar infarct (facio-brachial weakness, +/− dysphasia)
- Space-occupying lesions: intrinsic tumour, metastasis, abscess

**Bilateral:**
- Motor neurone disease
- Diffuse cerebrovascular disease
- Pontine infarct (locked-in syndrome)

- **Lower motor neurone facial weakness** (peripheral origin):
  If this is due to facial (VII) nerve palsy, it results in ipsilateral weakness of frontalis (cf. upper motor neurone facial paresis), orbicularis oculi, buccinator, orbicularis oris, and platysma. Clinically this produces
Facial Paresis, Facial Weakness

Drooping of the side of the face with loss of the nasolabial fold; widening of the palpebral fissure with failure of lid closure (lagophthalmos); eversion of the lower lid (ectropion) with excessive tearing (epiphora); inability to raise the eyebrow, close the eye, frown, blow out the cheek, show the teeth, laugh, and whistle; +/− dribbling of saliva from the paretic side of the mouth; depression of the corneal reflex (efferent limb of reflex arc affected); speech alterations: softening of labials (p, b).

Depending on the precise location of the facial nerve injury, there may also be paralysis of the stapedius muscle in the middle ear, causing sounds to seem abnormally loud (especially low tones: hyperacusis), and impairment of taste sensation on the anterior two-thirds of the tongue if the chorda tympani is affected (ageusia, hypogeusia). Lesions within the facial canal distal to the meatal segment cause both hyperacusis and ageusia; lesions in the facial canal between the nerve to stapedius and the chorda tympani cause ageusia but no hyperacusis; lesions distal to the chorda tympani cause neither ageusia nor hyperacusis (i.e. facial motor paralysis only). Lesions of the cerebellopontine angle cause ipsilateral hearing impairment and corneal reflex depression (afferent limb of reflex arc affected) in addition to facial weakness. There is also a sensory branch to the posterior wall of the external auditory canal which may be affected resulting in local hypoesthesia (Hitselberg sign).

Causes of lower motor neurone facial paresis include

• Bell’s palsy: idiopathic lower motor neurone facial weakness, assumed to result from a viral neuritis
• Herpes zoster (Ramsey Hunt syndrome)
• Diabetes mellitus
• Lyme disease (neuroborreliosis, Bannwarth’s disease)
• Sarcoïdosis
• Leukaemic infiltration, lymphoma
• HIV seroconversion
• Neoplastic compression (e.g. cerebellopontine angle tumour; rare)
• Facial nerve neuroma

These latter conditions may need to be differentiated from Bell’s palsy. Causes of recurrent facial paresis of lower motor neurone type include

• Diabetes mellitus
• Lyme disease (neuroborreliosis, Bannwarth’s disease)
• Sarcoïdosis
• Leukaemia, lymphoma

In myasthenia gravis, a disorder of neuromuscular transmission at the neuromuscular junction, there may be concurrent ptosis, diplopia, bulbar palsy, and limb weakness and evidence of fatiguable weakness. Myogenic facial paresis may be seen in facioscapulohumeral (FSH) dystrophy, myotonic dystrophy, and mitochondrial disorders. In primary disorders of muscle the pattern of weakness and family history may suggest the diagnosis.
References

Cross References
Abulia; Ageusia; Bell’s palsy; Bell’s phenomenon, Bell’s sign; Bouche de tapir; Cerebellopontine angle syndrome; Corneal reflex; Eight-and-a-half syndrome; Epiphora; Fisher’s sign; Hitselberg sign; Hyperacusis; Lagophthalmos; Locked-in syndrome; Lower motor neurone (LMN) syndrome; Pseudobulbar palsy; Upper motor neurone (UMN) syndrome

Facilitation
Facilitation is an increase in muscle strength following repeated contraction. Clinically, facilitation may be demonstrated by the appearance of tendon reflexes which are absent at rest after prolonged (ca. 30 s) forced maximal contractions against resistance, e.g. the biceps jerk after elbow flexion, knee jerk after knee extension; and by Lambert’s sign (increased force grip with sustained contraction; this increase in strength of affected muscles detected in the first few seconds of maximal voluntary contraction may also be known as augmentation).

This phenomenon of posttetanic potentiation is most commonly seen in the Lambert–Eaton myasthenic syndrome (LEMS), a disorder of neuromuscular junction transmission associated with the presence of autoantibodies directed against presynaptic voltage-gated calcium ion (Ca$^{2+}$) channels (VGCC). The mechanism is thought to be related to an increased build up of Ca$^{2+}$ ions within the presynaptic terminal with the repetitive firing of axonal action potentials, partially overcoming the VGCC antibody-mediated ion channel blockade, and leading to release of increasing quanta of acetylcholine.

Cross References
Augmentation; Fatigue; Lambert’s sign

False-Localizing Signs
Neurological signs may be described as ‘false-localizing’ when their appearance reflects pathology distant from the expected anatomical locus. The classic example, and probably the most frequently observed, is abducens nerve palsy (unilateral or bilateral) in the context of raised intracranial pressure, presumed to result from stretching of the nerve over the ridge of the petrous temporal bone. Many false-localizing signs occur in the clinical context of raised intracranial pressure, either idiopathic (idiopathic intracranial hypertension [IIH]) or symptomatic (secondary to tumour, haematoma, abscess). A brief topographical overview of false-localizing signs (more details may be found in specific entries) includes
• **Motor system:**
  - Kernohan’s notch syndrome: false-localizing hemiparesis
  - Cerebellar syndrome with anterior cerebral artery territory infarction damaging frontocerebellar pathways
  - Brainstem compression causing diaphragm paralysis

• **Sensory system:**
  - Sensory level with parietal lobe lesion

• **Cranial nerves:**
  - Proptosis with middle cranial fossa tumour
  - Oculomotor (III) nerve palsy with contralateral supratentorial lesion
  - Divisional oculomotor nerve palsy with brainstem or subarachnoid space pathology
  - Trochlear nerve palsy with IIH
  - Trigeminal nerve palsy with IIH
  - Abducens nerve palsy with IIH
  - Facial nerve palsy with IIH
  - Vestibulocochlear nerve dysfunction with IIH.

• **Spinal cord and roots:**
  - Foramen magnum/upper cervical cord lesion causing hand muscle wasting (‘remote atrophy’)
  - Lower cervical/upper thoracic myelopathy producing midthoracic girdle sensation
  - Urinary retention with rostral spinal cord compression
  - Radiculopathy with IIH, may even mimic Guillain–Barré syndrome

**References**


**Cross References**

Abducens (VI) nerve palsy; Divisional palsy; Girdle sensation; Kernohan’s notch syndrome; Oculomotor (III) nerve palsy; Proptosis; Urinary retention

**Fan Sign (Signe de L’éventail)**

- see BABINSKI’S SIGN (1)

**Fasciculation**

Fasciculations are rapid, flickering, twitching, involuntary movements within a muscle belly resulting from spontaneous activation of a bundle, or fasciculus, of muscle fibres (i.e. a motor unit), insufficient to move the joint. Fasciculations may also be induced by lightly tapping over a partially denervated muscle belly. The term was formerly used synonymously with fibrillation, but the latter term is now reserved for contraction of a single muscle fibre or a group of fibres smaller than a motor unit.

Brief and localized fasciculations can be a normal finding (e.g. in the intrinsic foot muscles, especially abductor hallucis, and gastrocnemius, but not tibialis anterior), particularly if unaccompanied by other neurological symptoms and
signs (wasting, weakness, sensory disturbance, sphincter dysfunction). Persistent fasciculations most usually reflect a pathological process involving the lower motor neurones in the anterior (ventral) horn of the spinal cord and/or in brain-stem motor nuclei, typically motor neurone disease (in which cramps are an early associated symptom). Facial and perioral fasciculations are highly characteristic of spinal and bulbar muscular atrophy (Kennedy’s disease). However, fasciculations are not pathognomonic of lower motor neurone pathology since they can on rare occasions be seen with upper motor neurone pathology.

The pathophysiological mechanism of fasciculations is thought to be spontaneous discharge from motor nerves, but the site of origin of this discharge is uncertain. Although ectopic neural discharge from anywhere along the lower motor neurone from cell body to nerve terminal could produce fasciculation, the commonly encountered assumption that this originates from the anterior horn cell body is not entirely supported by the available evidence, which points to an additional, more distal, origin in the motor axons. Denervation of muscle fibres may lead to nerve fibre sprouting (axonal and collateral) and enlargement of motor units which makes fasciculations more obvious clinically.

Fasciculations may be seen in:

- Motor neurone disease with lower motor neurone involvement (i.e. progressive muscular atrophy, progressive bulbar atrophy variants)
- Spinal muscular atrophy
- Cervical radiculopathy (restricted to myotomal distribution)
- Multifocal motor neuropathy with conduction block
- Benign fasciculation syndrome: typically seen only after exercise and without associated muscle atrophy or weakness
- Cramp fasciculation syndrome
- Spinal and bulbar muscular atrophy (Kennedy’s disease), especially perioral
- Almost any lower motor neurone disease, especially compression
- Metabolic causes: thyrotoxicosis, tetany, after acetylcholinesterase inhibitors, anaesthetic muscle relaxants

Fasciculations may need to be distinguished from myokymia or neuromyotonia.

**References**


**Cross References**

Calf hypertrophy; Cramp; Fibrillation; Lower motor neurone (LMN) syndrome; Myokymia; Neuromyotonia

**Fast Micrographia**

In ‘fast’ micrographia, written letters are microscopic from the outset, sometimes approximating to a straight line, though produced at normal speed without fatigue. This pattern has been observed in progressive supranuclear palsy and
with globus pallidus lesions, and contrasts with the ‘slow’ micrographia, meaning writing which becomes progressively slower and smaller, as seen in idiopathic Parkinson’s disease.

**Reference**

**Cross Reference**
Micrographia

**Fatigue**
The term fatigue may be used in different contexts to refer to both a sign and a symptom.

The sign of fatigue, also known as peripheral fatigue, consists of a reduction in muscle strength or endurance with repeated muscular contraction. This most characteristically occurs in disorders of neuromuscular junction transmission (e.g. myasthenia gravis), but it may also be observed in disorders of muscle (e.g. myopathy, polymyositis) and neurogenic atrophy (e.g. motor neurone disease). In myasthenia gravis, fatigue may be elicited in the extraocular muscles by prolonged upgaze causing eyelid drooping; in bulbar muscles by prolonged counting or speech causing hypophonia; and in limb muscles by repeated contraction, especially of proximal muscles (e.g. shoulder abduction, ‘wing flaps’) leading to weakness in previously strong muscles. Fatigue in myasthenia gravis is thought to be caused by a decline in the amount of acetylcholine released from motor nerve terminals with successive neural impulses, along with a reduced number of functional acetylcholine receptors (AChR) at the motor end-plates, due to binding of AChR antibodies and/or complement-mediated destruction of the postsynaptic folds.

A gradual decline in the amplitude and speed of initiation of voluntary movements, hypometria and hypokinesia, as seen in disorders of the basal ganglia, especially Parkinson’s disease, may also be described as fatigue, e.g. ‘slow’ micrographia may be ascribed to ‘fatigue’. Progressive supranuclear palsy is notable for lack of such fatigue.

Fatigue as a symptom, or central fatigue, is an enhanced perception of effort and limited endurance in sustained physical and mental activities. This may occur in multiple sclerosis (MS), post-polio syndrome, post-stroke syndromes, and chronic fatigue syndrome (CFS). In MS and CFS, fatigue may be a prominent and disabling complaint even though neurological examination reveals little or no clinical deficit. This type of fatigue is ill-understood: in MS, frequency-dependent conduction block in demyelinated axons has been suggested, as has hypothalamic pathology. Current treatment is symptomatic (amantadine, modafanil, 3,4-diaminopyridine) and rehabilitative (graded exercise).

Fatigue may be evaluated with various instruments, such as the Krupp Fatigue Severity Score.

**References**
Cross References
Dystonia; Hypokinesia; Hypometria; Micrographia; Weakness

Femoral Stretch Test
The femoral stretch test, or reverse straight leg raising test, consists of extension of the hip with the knee straight with the patient lying prone, a manoeuvre which puts traction on the femoral nerve or L3 root and may exacerbate pain in a femoral neuropathy or L3 radiculopathy, perhaps caused by a retroperitoneal haemorrhage.

Cross Reference
Lasègue’s sign

Fencer’s Posture, Fencing Posture
Epileptic seizures arising in or involving the supplementary motor area may lead to adversial head and eye deviation, abduction and external rotation of the contralateral arm, flexion at the elbows, and posturing of the legs, with maintained consciousness, a phenomenon christened by Penfield as the ‘fencing posture’ because of its resemblance to the en garde position. These may also be known as ‘salutatory seizures’.

Cross Reference
Seizures

Festinant Gait, Festination
Festinant gait or festination is a gait disorder characterized by rapid short steps (Latin: festinare, to hurry, hasten, accelerate) due to inadequate maintenance of the body’s centre of gravity over the legs. To avoid falling and to maintain balance the patient must ‘chase’ the centre of gravity, leading to an increasing speed of gait and a tendency to fall forward when walking (propulsion). A similar phenomenon may be observed if the patient is pulled backwards (retropulsion). Festination may be associated with freezing of gait.

Festation is common in idiopathic Parkinson’s disease; it is associated with longer duration of disease and higher Hoehn & Yahr stage. Festation may be related to the flexed posture and impaired postural reflexes commonly seen in these patients. It is less common in symptomatic causes of parkinsonism, but has been reported, for example, in aqueduct stenosis.

Reference

Cross References
Freezing; Parkinsonism; Postural reflexes

Fibrillation
Fibrillation was previously synonymous with fasciculation, but the term is now reserved for the spontaneous contraction of a single muscle fibre, or a group of fibres smaller than a motor unit, hence this is more appropriately regarded as an electrophysiological sign without clinical correlate.
Finger Agnosia

Finger agnosia is a type of tactile agnosia, in which there is inability to identify which finger has been touched when the eyes are closed, despite knowing that a finger has been touched; or inability to point to or move a finger when it is named; or inability to name the fingers (patient’s own fingers or those of another person). This is a disorder of body schema and may be regarded as a partial form of autotopagnosia.

Finger agnosia is most commonly observed with lesions of the dominant parietal lobe. It may occur in association with agraphia, agraphia, and right–left disorientation, with or without agraphia and difficulty spelling words, hence as one feature of Gerstmann syndrome. Isolated cases of finger agnosia in association with left cortic-subcortical posterior parietal infarction have been reported. Since this causes no functional deficit, it may be more common than reported.

Reference

Cross References
Agnosia; Autotopagnosia; Gerstmann syndrome

Finger Chase Test
- see ATAXIA; CEREBELLAR SYNDROMES

Finger Drop
- see WRIST DROP

Finger–Floor Distance
In patients with leg (+/-low back) pain suspected of having lumbosacral nerve root compression, a finger-floor distance of >25 cm when the patient bends forward and attempts to touch the floor with the fingers has been found to be an independent predictor of radiological (MR imaging) compression. This was not the case for the straight leg raising test.

Reference

Cross Reference
Lasègue’s sign

Finger-to-Nose Test
- see ATAXIA; CEREBELLAR SYNDROMES
**Fisher’s Sign**
Fisher’s sign is the paucity of facial expression conveying emotional states or attitudes (emotional facial paresis). It follows non-dominant (right) hemisphere lesions and may accompany emotional dysprosody of speech.

**Cross References**
Abulia; Aprosodia, Aprosody; Facial paresis, Facial weakness

**Fist-Edge-Palm Test**
In the fist-edge-palm test, sometimes known as the Luria test or three-step motor sequence, the patient is requested to place the hand successively in three positions, imitating movements made by the examiner and then doing them alone: fist, vertical palm, palm resting flat on table. Copying motor sequences assesses motor programming ability. Defects in this programming, such as lack of kinetic melody, loss of sequence, or repetition of previous pose or position, are especially conspicuous with anterior cortical lesions. This test is incorporated into the Frontal Assessment Battery.

**Reference**

**Cross Reference**
Frontal lobe syndromes

**Flaccidity**
Flaccidity is a floppiness which implies a loss of normal muscular tone (hypotonia). This may occur transiently after acute lesions of the corticospinal tracts (flaccid paraparesis), before the development of spasticity, or as a result of lower motor neurone syndromes. It is difficult to separate the change in tone from weakness.

**Cross References**
Hypotonia, Hypotonus; Lower motor neurone (LMN) syndrome

**Flail Arm**
Flail arm refers to a severe and symmetric wasting and weakness of the arms without significant functional involvement of other regions, seen in one variant of motor neurone disease, the ‘flail arm syndrome’, also known as Vulpian–Bernhart’s form. Men are reported to be much more frequently affected than women, and this group may show improved survival compared to other MND patients. Alternative designations for this syndrome include amyotrophic brachial diplegia, dangling arm syndrome, and neurogenic man-in-a-barrel syndrome. It has been suggested that prognosis may be relatively good in this variant as compared to other manifestations of MND. Cognition may be relatively well preserved.

**References**


**Cross References**

Amyotrophy; ‘Man-in-a-barrel’

**Flail Foot**

- see CAUDA EQUINA SYNDROME; FOOT DROP

**Flap, Flapping Tremor**

- see ASTERIXIS

**Flexibilitas Cerea (Waxy Flexibility)**

- see CATATONIA

**Flexion–Adduction Sign**

Neuralgic amyotrophy (Parsonage–Turner syndrome) may cause arm pain, which may be prevented by holding the arm flexed at the elbow and adducted at the shoulder.

**Reference**


**Flexor Posturing**

- see DECORTICATE RIGIDITY

**Flick Sign**

A flicking, shaking movement of the hands made by patients with carpal tunnel syndrome to try to relieve the paraesthesia and pain caused by the condition, typically noted on waking at night. This may be the most sensitive and specific of the various signs described in carpal tunnel syndrome.

**References**


**Cross References**

Phalen’s sign; Tinel’s sign

**Floccillation**

- see CARPHOLOGIA

**Flycatcher Tongue**

- see TROMBONE TONGUE
**Flynn Phenomenon**
Flynn phenomenon is paradoxical constriction of the pupils in darkness. This has been documented in various conditions including congenital achromatopsia, following optic neuritis, and in autosomal dominant optic atrophy.

**Reference**

**Cross Reference**
Pupillary reflexes

**Foot Drop**
Foot drop, often manifest as the foot dragging during the swing phase of the gait, causing tripping and/or falls, may be due to upper or lower motor neurone lesions, which may be distinguished clinically.

- **Stiff foot drop**, with upper motor neurone lesions:
  leads to a circumducting gait; it may be possible to see or hear the foot dragging or scuffing along the floor, and this may cause excessive wear on the point of the shoe. There will be other upper motor neurone signs (hemiparesis; spasticity, clonus, hyperreflexia, Babinski’s sign).

- **Floppy foot drop**, with lower motor neurone lesions:
  leads to a stepping gait (steppage) to try to lift the foot clear of the floor in the swing phase, and a slapping sound on planting the foot. At worst, there is a flail foot in which both the dorsiflexors and the plantar flexors of the foot are weak (e.g. in high sciatic nerve or sacral plexus lesions). Other lower motor neurone signs may be present (hypotonia, areflexia, or hyporeflexia).

Causes of floppy foot drop include

- Common peroneal nerve palsy
- Sciatic neuropathy
- Lumbosacral plexopathy
- L4/L5 radiculopathy
- Motor or sensorimotor polyneuropathy (e.g. hereditary motor and sensory neuropathy)
- Motor neuronopathy (anterior horn cell disease)
- Mononeuropathy multiplex

These may be distinguished on clinical and/or neurophysiological grounds

**Reference**

**Cross References**
Cauda equina syndrome; Hemiparesis; Lower motor neurone (LMN) syndrome; Steppage, Stepping gait; Upper motor neurone (UMN) syndrome

**Foot Grasping**
- see GRASP REFLEX
**Forced Ductions**
Forced ductions, performed by grasping the anaesthetized sclera with forceps and then moving the eye through its range of motions, may be used to determine whether restricted eye movement is mechanical, due to a lesion within the orbit, such as thyroid ophthalmopathy or superior oblique tendon sheath (Brown’s) syndrome.

**Forced Grasping**
- see GRASP REFLEX

**Forced Groping**
Forced groping describes involuntary movements of a hand, as if searching for an object or item which has touched or brushed against it; the hand may follow the object around if it moves (magnetic movements). There may be an accompanying grasp reflex. This type of behaviour may be displayed by an alien hand, most usually in the context of corticobasal degeneration. Forced groping may be conceptualized as an exploratory reflex which is ‘released’ from frontal lobe control by a pathological process, as in utilization behaviour.

Reference

Cross References
Alien hand, Alien limb; Grasp reflex; Magnetic movements; Utilization behaviour

**Forced Laughter and Crying**
- see EMOTIONALISM, EMOTIONAL LABILITY; PATHOLOGICAL CRYING, PATHOLOGICAL LAUGHTER

**Forced Upgaze**
Tonic upward gaze deviation, forced upgaze, may be seen in coma after diffuse hypoxic–ischaemic brain injury with relative sparing of the brainstem. Forced upgaze may also be psychogenic, in which case it is overcome by cold caloric stimulation of the ear drums. Forced upgaze must be differentiated clinically from oculogyric crisis.

Cross Reference
Oculogyric crisis

**Forearm and Finger Rolling**
The forearm and finger rolling tests detect subtle upper motor neurone lesions with high specificity and modest sensitivity. Either the forearms or the index fingers are rapidly rotated around each other in front of the torso for about 5 s, then the direction reversed. Normally the appearance is symmetrical but with a unilateral upper motor neurone lesion one arm or finger remains relatively stationary, with the normal rotating around the abnormal limb. Thumb rolling might also be a sensitive test for subtle upper motor neurone pathology.

Reference
Cross References
Pronator drift; Upper motor neurone (UMN) syndrome

Foreign Accent Syndrome
The foreign accent syndrome is a rare phonological disorder, such that speech production includes non-native vowels or consonants and hence sounds as though it is foreign, or different from the speakers native intonation. There is no language disorder since comprehension of spoken and written language is preserved; hence it is qualitatively different from Broca's aphasia. This syndrome probably overlaps with other disorders of speech production, labelled as phonetic disintegration, pure anarthria, aphemia, apraxic dysarthria, verbal or speech apraxia, and cortical dysarthria. Case heterogeneity is noted; some cases may be non-organic.

References

Cross References
Aphasia; Aphemia

Formication
Formication is a tactile hallucination, as of ants crawling over the skin.

Cross References
Hallucination; Paraesthesia; Tinel's sign

Fortification Spectra
Fortification spectra, also known as teichopsia, are visual hallucinations which occur as an aura, either in isolation (migraine aura without headache) or prior to an attack of migraine (migraine with aura; ‘classical migraine’). The appearance is a radial array likened to the design of medieval castles, not simply of battlements. Hence these are more complex visual phenomena than simple flashes of light (photopsia) or scintillations. They are thought to result from spreading depression, of possible ischaemic origin, in the occipital cortex. The visions of Hildegard von Bingen (1098–1179), illustrated in the twelfth century, are thought possibly to reflect migrainous fortification spectra.

Reference

Cross References
Aura; Hallucination; Photopsia; Teichopsia

Foster Kennedy Syndrome
The Foster Kennedy syndrome consists of optic atrophy in one eye with optic disc oedema in the other eye, Anosmia ipsilateral to optic atrophy may also be found.
This is classically due to a tumour, typically an olfactory groove meningioma, which compresses the ipsilateral optic nerve to cause atrophy, and raises intracranial pressure to cause contralateral papilloedema. Similar clinical appearances may occur with sequential anterior ischaemic optic neuropathy, sometimes called a pseudo-Foster Kennedy syndrome.

Reference
Kennedy F. Retrobulbar neuritis as an exact diagnostic sign of certain tumors and abscesses in the frontal lobe. *American Journal of Medical Science* 1911; **142**: 355–368.

Cross References
Optic atrophy; Papilloedema

**Fou Rire Prodromique**

*Fou rire prodromique*, or laughing madness, first described by Féré in 1903, is pathological laughter which heralds the development of a brainstem stroke, usually as a consequence of basilar artery occlusion. Pathological crying as a prodrome of brainstem stroke has also been described (‘folles larmes prodromiques’).

References


**Freezing**

Freezing is the sudden inability in a patient with parkinsonism to move or to walk, i.e. gait failure, as though the patient were turned to ice or the feet were nailed to the floor. This is one of the unpredictable motor fluctuations in late Parkinson’s disease (associated with longer duration of disease and treatment) which may lead to falls, usually forward onto the knees, and injury. It may occur in confined spaces (e.g. doorways), when trying to turn, or when trying to do two things at once. It is not seen in the early years of levodopa therapy. Two variants are encountered, occurring either during an off period or wearing off period, or randomly, i.e. unrelated to drug dosage or timing.

Treatment strategies include use of dopaminergic agents and, anecdotally, L-threodops, but these agents are not reliably helpful, particularly in random freezing. Use of visual targets (real or imagined) may help, e.g. stepping over a line. Freezing may also occur in multiple system atrophy and has also been reported as an isolated phenomenon.

Reference

Cross Reference
Parkinsonism

**Fregoli Syndrome**

- see DELUSION
Frey’s Syndrome
- see GUSTATORY SWEATING

Froment’s Sign
Froment has two eponymous signs:
- Activated rigidity or synkinesis; sometimes known as Froment’s manoeuvre.
- In an ulnar nerve lesion, flexion of the distal phalanx of the thumb (flexor pollicis longus, innervated by the median nerve) is seen when attempting to squeeze a sheet of paper between the thumb and the index finger, as a compensation for the weakness of thumb adduction (adductor pollicis, innervated by the ulnar nerve), also known as Froment’s prehensile thumb sign or the signe du journal. The term is also sometimes used for weakness of little finger adduction (palmar interossei), evident when trying to grip a piece of paper between the ring and little finger.

Cross References
Rigidity; Synkinesia, Synkinesis; Wartenberg’s sign (1)

Frontal Ataxia
- see ATAXIA

Frontal Lobe Syndromes
The frontal lobes of the brain have enlarged greatly during phylogeny; their diverse connections with the basal ganglia, basal forebrain, and cerebellum, as well as other cortical areas, reflect their multiple motor and behavioural functions. Damage to the frontal lobes may produce a variety of clinical signs, most frequently changes in behaviour. Such changes may easily be overlooked with the traditional neurological examination, although complained of by patient’s relatives, and hence specific bedside tests of frontal lobe function should be utilized, for example:
- Verbal fluency: e.g. letter/phonemic (F, A, S) probably a more specific test than category/semantic (animals, foods).
- Proverb interpretation: e.g. ‘Make hay while the sun shines’ ‘Too many cooks spoil the broth’; interpretation tends to be concrete in frontal lobe disorders.
- Cognitive estimates: e.g. height of the Post Office Tower, length of a man’s spine, distance from London to Edinburgh; may be grossly abnormal or inappropriate.
- Copying motor sequences, to assess motor programming ability: e.g. Luria fist-edge-palm test (three-step motor sequence with hand).
- Alternating sequence tests: e.g. alternating finger flexion/extension out of phase in two hands, or repeatedly writing m n m n m n (also used as tests of praxis, which may be affected with frontal lobe pathology); swapping a coin from hand to hand behind back in a predictable pattern and asking the patient which hand the coin is in.
- Set-shifting or go/no-go tests, in which an alternating pattern is suddenly changed, e.g. changing the previously predictable (left/right) pattern of coin hidden in clenched hand swapped over behind back; rhythmic tapping with pen on a surface (I tap once, you tap twice; I tap twice, you tap once); tests
of response inhibition (ask patient to clap three times, s/he does so multiple times).

A useful clinico-anatomical classification of frontal lobe syndromes which reflects the functional subdivisions of the frontal lobes is as follows:

- **Orbitofrontal syndrome (‘disinhibited’):**
  - disinhibited behaviour (including sexual disinhibition), impulsivity
  - inappropriate affect, *witzelsucht*, euphoria
  - emotional lability (moria)
  - lack of judgement, insight
  - distractibility, lack of sustained attention; hypermetamorphosis
  - motor perseverations are not a striking feature

- **Frontal convexity syndrome (‘apathetic’):**
  - apathy; abulia, indifference
  - motor perseveration
  - difficulty set-shifting, stimulus boundedness
  - reduced verbal fluency
  - deficient motor programming, e.g. three-step hand sequence, rhythmic tapping (go/no-go test).

- **Medial frontal syndrome (‘akinetic’):**
  - little spontaneous movement, bradykinesia, hypokinesia
  - sparse verbal output (akinetic mutism)
  - urinary incontinence
  - sensorimotor signs in lower limbs
  - indifference to pain

Overlap between these regional syndromes may occur.

A ‘dysexecutive syndrome’ has also been defined, consisting of difficulty planning, adapting to changing environmental demands (impaired cognitive flexibility, e.g. in set-shifting tests), and directing attentional resources. This may be seen with dorsolateral (prefrontal) damage.

These frontal lobe syndromes may be accompanied by various neurological signs (frontal release signs or primitive reflexes). Other phenomena associated with frontal lobe pathology include imitation behaviours (echophenomena) and, less frequently, utilization behaviour, features of the environmental dependency syndrome.

Frontal lobe syndromes may occur as a consequence of various pathologies:

- Neurodegenerative diseases: especially frontal or behavioural variant frontotemporal lobar degeneration; occasionally in Alzheimer’s disease;
- Structural lesion: tumour (intrinsic, extrinsic), normal pressure hydrocephalus;
- Cerebrovascular event;
- Head injury;
- Inflammatory metabolic disease: multiple sclerosis, X-linked adrenoleucodystrophy.
Frontal Release Signs

Frontal release signs are so named because of the belief that they are released from frontal inhibition by diffuse pathology within the frontal lobes (usually vascular or degenerative) with which they are often associated, although they may be a feature of normal ageing. Some of these responses are present during infancy but disappear during childhood, hence the terms ‘primitive reflexes’ or ‘developmental signs’ are also used (Babinski’s sign may therefore fall into this category). The term ‘psychomotor signs’ has also been used since there is often accompanying change in mental status. The frontal release signs may be categorized as:

- **Prehensile:**
  - Sucking reflex (tactile, visual)
  - Grasp reflex: hand, foot
  - Rooting reflex (turning of the head towards a tactile stimulus on the face)

- **Nociceptive:**
  - Snout reflex
  - Pout reflex
  - Glabellar (blink) reflex
  - Palmomental reflex

The corneomandibular and nucrocephalic reflexes may also be categorized as ‘frontal release’ signs.

Some are of little clinical value (e.g. palmomental reflex). Concurrent clinical findings may include dementia, gait disorder (frontal gait, *marche à petit pas*), urinary incontinence, akinetic mutism, and *gegenhalten*. Common causes of these findings are diffuse cerebrovascular disease and motor neurone disease, and they...
may be more common in dementia with Lewy bodies than other causes of an extrapyramidal syndrome. All increase with age in normal individuals.

References

Cross References
Age-related signs; Babinski’s sign (1); Corneomandibular reflex; *Gegenhalten*; Grasp reflex; *Marche à petit pas*; Palmomental reflex; Pout reflex; Rooting reflex; Sucking reflex

**Fugue**
Fugue, and fugue-like state, is used to refer to a syndrome characterized by loss of personal memory (hence the alternative name of ‘twilight state’), automatic and sometimes repetitive behaviours, and wandering or driving away from normal surroundings. Fugue may be:
- Psychogenic: associated with depression (sometimes with suicide); alcoholism, amnesia; ‘hysteria’
- Epileptic: complex partial seizures
- Narcoleptic

Some patients with frontotemporal dementia may spend the day walking long distances, and may be found a long way from home, unable to give an account of themselves, and aggressive if challenged; generally they are able to find their way home (spared topographical memory) despite their other cognitive deficits.

Cross References
Amnesia; Automatism; Dementia; Poriomania; Seizures

**Functional Weakness and Sensory Disturbance**
Various signs have been deemed useful indicators of functional or ‘non-organic’ neurological illness, including
- Collapsing or ‘give way’ weakness
- Hoover’s sign
- Babinski’s trunk–thigh test
- ‘Arm drop’
- *Belle indifference*
- Sternocleidomastoid sign
- Midline splitting sensory loss
- Functional postures, gaits:
  - monoplegic ‘dragging’
  - fluctuation of impairment
excessive slowness, hesitation
‘psychogenic Romberg’ sign
‘walking on ice’
uneconomic posture, waste of muscle energy.
sudden knee buckling

Although such signs may be suggestive, their diagnostic utility has never been formally investigated in prospective studies, and many, if not all, have been reported with ‘organic’ illness. Hence it is unwise to rely on them as diagnostic indicators.

References

Cross References
‘Arm drop’; Babinski’s trunk–thigh test; *Belle indifférence*; Collapsing weakness; Harvey’s sign; Hawthorne effect; Hoover’s sign; Sternocleidomastoid test

Funnel Vision
- see ‘TUNNEL VISION’
Gag Reflex
The gag reflex is elicited by touching the posterior pharyngeal wall, tonsillar area, or the base of the tongue, with the tip of a thin wooden (‘orange’) stick. Depressing the tongue with a wooden spatula, and the use of a torch for illumination of the posterior pharynx, may be required to get a good view. There is a palatal response (palatal reflex), consisting of upward movement of the soft palate with ipsilateral deviation of the uvula; and a pharyngeal response (pharyngeal reflex or gag reflex) consisting of visible contraction of the pharyngeal wall. Lesser responses include medial movement, tensing, or corrugation of the pharyngeal wall. In addition there may be head withdrawal, eye watering, coughing, and retching. Hence there is variability of response in different individuals. Some studies claim that the reflex is absent in many normal individuals, especially with increasing age, without evident functional impairment; whereas others find it in all healthy individuals, although variable stimulus intensity is required to elicit it.

The afferent limb of the reflex arc is the glossopharyngeal (IX) nerve, the efferent limb is the glossopharyngeal and vagus (X) nerves. Hence individual or combined lesions of the glossopharyngeal and vagus nerves depress the gag reflex, as in neurogenic bulbar palsy.

Dysphagia is common after a stroke, and the gag reflex is often performed to assess the integrity of swallowing. Some argue that absence of the reflex does not predict aspiration and is of little diagnostic value, since this may be a normal finding in elderly individuals, whereas pharyngeal sensation (feeling the stimulus at the back of the pharynx) is rarely absent in normals and is a better predictor of the absence of aspiration. Others find that even a brisk pharyngeal response in motor neurone disease may be associated with impaired swallowing. Hence the value of the gag reflex remains debatable. A video swallow may be a better technique to assess the integrity of swallowing.

References

Cross References
Bulbar palsy; Dysphagia

Gait Apraxia
Gait apraxia is a name given to an inability to walk despite intact motor systems and sensorium. Patients with gait apraxia are often hesitant, seemingly unable to lift their feet from the floor (‘magnetic gait’) or put one foot in front of the other.
Arms may be held out at the sides to balance for fear of falling; fear may be so great that the patient sits in a chair gripping its sides. These phenomena may be observed with lesions of the frontal lobe and white matter connections, with or without basal ganglia involvement, for example, in diffuse cerebrovascular disease and normal pressure hydrocephalus. A syndrome of isolated gait apraxia has been described with focal degeneration of the medial frontal lobes. In modern classifications of gait disorders, gait apraxia is subsumed into the categories of frontal gait disorder, frontal disequilibrium, and isolated gait ignition failure.

Gait apraxia is an important diagnosis to establish since those afflicted generally respond poorly, if at all, to physiotherapy; moreover, because both patient and therapist often become frustrated because of lack of progress, this form of treatment is often best avoided.

References

Cross Reference
Apraxia

**Gambling**
Gambling may be characterized as an executive function task, amenable to testing with instruments such as the Iowa Gambling Task (IGT) and the Cambridge Gamble Task. The neuroanatomical substrates of such decision-making are believed to encompass the prefrontal cortex and the amygdala. Gambling may be defined as pathological when greater risks are taken and potential losses are correspondingly greater; this may be classified as an impulse control disorder. Pathological gambling may occur in patients with Parkinson’s disease treated with various dopamine agonists and in frontal variant frontotemporal dementia patients who display risky decision-making, even in early disease and without evidence of behavioural disinhibition or impulsiveness.

References

Cross Reference
Punding

**Ganglionopathy**
- see NEUROPATHY

**Ganser Phenomenon, Ganser Syndrome**
The Ganser phenomenon consists of giving approximate answers to questions which can at times verge on the absurd (Q: ‘How many legs does a cow have?’; A: ‘Three’), also known as paralogia or *vorbereiden*. This may occur in psychiatric
disease such as depression, schizophrenia, and malingering, and sometimes in neurological disease (head injury, epilepsy). A Ganser syndrome of hallucinations, conversion disorder, cognitive disorientation, and approximate answers is also described but of uncertain nosology.

References

Gaping
Gaping, or involuntary opening of the mouth, may occur as a focal dystonia of the motor trigeminal nerve, also known as Brueghel syndrome after that artist’s painting De Gaper (‘Yawning man’, ca. 1558) which is said to illustrate a typical case. Afflicted individuals may also demonstrate paroxysmal hyperpnoea and upbeat nystagmus, suggesting a brainstem (possibly pontine) localization of pathology. The condition should be distinguished from other cranial dystonias with blepharospasm (Meige syndrome).

Reference

Cross References
Blepharospasm; Dystonia; Nystagmus

Gaze-Evoked Phenomena
A variety of symptoms have been reported to be evoked, on occasion, by alteration of the direction of gaze:

- **Amaurosis**: lesion, usually intraorbital, compressing central retinal artery;
- **Laughter**;
- **Nystagmus**: usually indicative of a cerebellar lesion; may occur as a side-effect of medications; also convergence–retraction nystagmus on upgaze in dorsal midbrain (Parinaud’s) syndrome;
- **Phosphenes**: increased mechanosensitivity in demyelinated optic nerve;
- **Segmental constriction of the pupil** (Czarnecki’s sign) following aberrant regeneration of the oculomotor (III) nerve to the iris sphincter;
- **Tinnitus**: may develop after resection of cerebellopontine angle tumours, may be due to abnormal interaction between vestibular and cochlear nuclei;
- **Vertigo**.

Reference

Gaze Palsy
Gaze palsy is a general term for any impairment or limitation in conjugate (yoked) eye movements. This may be supranuclear, nuclear, or infranuclear in origin. Preservation of the vestibulo-ocular reflexes may help differentiate supranuclear gaze palsies from nuclear/infranuclear causes.
Cross References
Locked-in syndrome; Supranuclear gaze palsy; Vestibulo-ocular reflexes

**Gegenhalten**

_Gegenhalten_, or paratonia, or paratonic rigidity, is a variable resistance to passive movement of a limb when changing its posture or position, which is evident in both flexor and extensor muscles (as in rigidity, but not spasticity), which seems to increase further with attempts to get the patient to relax, such that there is a resistance to any applied movement (German: to counter, stand ones ground). However, this is not a form of impaired muscle relaxation akin to myotonia and paramyotonia. For instance, when lifting the legs by placing the hands under the knees, the legs may be held extended at the knees despite encouragement on the part of the examiner for the patient to flex the knees. Generally, tendon reflexes are normal, plantar responses downgoing, and there is no clonus. _Gegenhalten_ is a sign of bilateral frontal lobe dysfunction, especially mesial cortex and superior convexity (premotor cortex, area 6). It is not uncommon in otherwise healthy elderly individuals with diffuse frontal lobe cerebrovascular disease.

Cross References
Frontal release signs; Myotonia; Paramyotonia; Rigidity; Spasticity

**Geophagia, Geophagy**

Geophagia or geophagy describes earth or clay eating, reports of which dating back to Hippocrates have been found. This may also fall under the rubric of pica, or pagophagia, a morbid craving for unusual or unsuitable food. Besides the obvious risk of infection from ingesting potentially contaminated material, geophagia may be associated with neurological complications. Cases of flaccid quadriplegia and of proximal myopathy associated with profound hypokalaemia in the context of geophagia have been reported, which may lead to walking difficulty.

References

**Gerstmann Syndrome**

The Gerstmann syndrome, or angular gyrus syndrome, consists of acalculia, agraphia (of central type), finger agnosia, and right–left disorientation; there may in addition be alexia and difficulty spelling words but these are not necessary parts of the syndrome.

Gerstmann syndrome occurs with lesions of the angular gyrus and supramarginal gyrus in the posterior parietotemporal region of the dominant (usually left) hemisphere, for example, infarction in the territory of the middle cerebral artery.

All the signs comprising Gerstmann syndrome do fractionate or dissociate, i.e. they are not causally related, or representative of a unitary neuropsychological function, as was once suggested. Hence this may be an example of a
disconnection syndrome. Nonetheless the Gerstmann syndrome remains useful for the purposes of clinical localization.

**References**


**Cross References**

Acalculia; Agraphia; Alexia; Finger agnosia; Right–left disorientation

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**Geste Antagoniste**

*Geste antagoniste* is a sensory ‘trick’ which alleviates, and is characteristic of, dystonia. *Geste antagoniste* consists of a tactile or proprioceptive stimulus, which is learned by the patient, which reduces or eliminates the dystonic posture. For example, touching the chin, face, or neck may overcome cervical dystonia (torticollis), and singing may inhibit blepharospasm. *Gestes* may also modify tremor. They are almost ubiquitous in sufferers of cervical dystonia and have remarkable efficacy. The *geste* phenomenon is said to be absent in psychogenic dystonia.

The mechanism is unknown: although afferent feedback from the periphery may be relevant, it is also possible that concurrent motor output to generate the trick movement may be the key element, in which case the term ‘sensory trick’ is a misnomer.

**References**


**Cross References**

Dystonia; Reverse sensory geste; Torticollis

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**Gibbus**

Angulation of the spine due to vertebral collapse may be due to osteoporosis, metastatic disease, or spinal tuberculosis. There may be associated myelopathy. Camptocormia (bent spine syndrome) enters the differential diagnosis.

**Cross References**

Camptocormia; Myelopathy

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**Girdle Sensation**

Compressive lower cervical or upper thoracic myelopathy may produce spastic paraparesis with a false-localizing midthoracic sensory level or ‘girdle sensation’ (cf. cuirasse). The pathophysiology is uncertain, but ischaemia of the thoracic
watershed zone of the anterior spinal artery from compression at the cervical level has been suggested.

**Reference**

**Cross References**
‘False-localizing signs’; Paraparesis; Suspended sensory loss

‘Give-Way’ Weakness - see COLLAPSING WEAKNESS; FUNCTIONAL WEAKNESS AND SENSORY DISTURBANCE

**Glabellar Tap Reflex**
The glabellar tap reflex, also known as Myerson’s sign or the nasopalpebral reflex, is elicited by repeated gentle tapping with a finger on the forehead, preferably with irregular cadence and so that the patient cannot see the finger (to avoid blinking due to the threat or menace reflex), whilst observing the eyelids blink (i.e. blink reflex). Usually, reflexive blinking in response to tapping habituates quickly, but in extrapyramidal disorders it may not do so. This sign was once thought useful for the diagnosis of idiopathic Parkinson’s disease but in fact it is fairly non-specific, occurring in many akinetic-rigid disorders.

**References**

**Cross References**
Blink reflex; Parkinsonism

**Glossolalia**
Glossolalia, or speaking in tongues, may be considered a normal phenomenon in certain Christian denominations, as divinely inspired, since it is mentioned in the Bible (1 Corinthians, 14:27–33, although St Paul speaks of the importance of an interpreter, since ‘God is not the author of confusion’), but it is not confined to Christianity or even overtly religious environments. Others conceptualize glossolalia as a form of automatic speech, usually of a pseudolanguage which may be mistaken for a foreign tongue. Such happenings may occur in trance-like states or in pathological states such as schizophrenia.

**Reference**

‘Glove and Stocking’ Sensory Loss
Sensory loss, to all or selected modalities, confined to the distal parts of the limbs (‘glove and stocking’) implies the presence of a peripheral sensory neuropathy.
If the neuropathy involves both sensory and motor fibres, motor signs (distal weakness, reflex diminution or loss) may also be present.

**Cross Reference**
Neuropathy

**Goosebumps**
- see ANSERINA

**Gordon’s Sign**
Gordon’s sign is an extensor plantar response in response to squeezing the calf muscles, also called the paradoxical flexor response. As with Chaddock’s sign and Oppenheim’s sign, this reflects an expansion of the receptive field of the reflex.

**Cross References**
Babinski’s sign (1); Plantar response

**Gowers’ Manoeuvre, Gowers’ Sign**
Gowers’ sign is a characteristic manoeuvre used by patients with proximal lower limb and trunk weakness to rise from the ground. From the lying position, the patient rolls to the kneeling position, pushes on the ground with extended forearms to lift the hips and straighten the legs, so forming a triangle with the hips at the apex with hands and feet on the floor forming the base (known in North America as the ‘butt-first manoeuvre’). Then the hands are used to push on the knees and so lift up the trunk (‘climbing up oneself’).

This sign was originally described by Gowers in the context of Duchenne muscular dystrophy but may be seen in other causes of proximal leg and trunk weakness, e.g. Becker muscular dystrophy, spinal muscular atrophy. Gowers was not the first to describe the sign; Bell had reported it almost 50 years before Gowers’ account.

Gowers’ name is also associated with a manoeuvre to stretch the sciatic nerve and hence exacerbate sciatic symptoms.

**Reference**

**Graefe’s Sign**
- see VON GRAEFE’S SIGN

**Grandchild Sign**
This sign is present if the child of an individual with dementia answers in the negative when asked if they would allow the patient to drive their own children (i.e. the patient’s grandchildren). It is said to be an indicator that the demented patient should not be driving.

**Reference**

**Graphaesthesia**
Graphaesthesia is the ability to identify numbers or letters written or traced on the skin, first described by Head in 1920. Loss of this ability (agraphaesthesia,
dysgraphaesthesia, or graphanaesthesia; sometimes referred to as agraphagnosia) is typically observed with parietal lobe lesions, for example, in conditions such as corticobasal degeneration. Such a cortical sensory syndrome may also cause astereognosis and impaired two-point discrimination.

Cross References
Agraphaesthesia; Astereognosis; Two-point discrimination

Graphanaesthesia
- see AGRAPHAESTHESIA

Graphospasm
- see WRITER’S CRAMP

Grasp Reflex
The grasp reflex consists of progressive forced closure of the hand (contraction of flexor and adductor muscles) when tactile stimulation (e.g. the examiner’s hand) is moved slowly, exerting pressure, across the patient’s palm in an upward direction. Once established, the patient is unable to release the grip (forced grasping), allowing the examiner to draw the arm away from the patient’s body. There may also be accompanying groping movements of the hand, once touched, in search of the examiner’s hand or clothing (forced groping, magnetic movement). Although categorized as a reflex, it may sometimes be accessible to modification by will (so-called alien grasp reflex). It is usually bilateral, even with unilateral pathology. Foot grasping (i.e. flexion and adduction of the toes and curling of the sole in response to pressure on the sole) may coexist, as may other frontal release signs (e.g. pout reflex, palmomental reflex, gegenhalten).

The grasp reflex may be categorized as a frontal release sign (or primitive reflex) of prehensile type, since it is most commonly associated with lesion(s) in the frontal lobes or deep nuclei and subcortical white matter. Clinicoradiological correlations suggest that the cingulate gyrus is the structure most commonly involved, followed by the supplementary motor area. Luria maintained that forced grasping resulted from extensive lesions of premotor region, disturbing normal relationships with the basal ganglia.

References

Cross References
Akinetic mutism; Alien grasp reflex; Frontal release signs

Groucho Marx Manoeuvre
Named for the American comic actor Julius Henry ‘Groucho’ Marx (1890–1977), this manoeuvre requires the forehead to be wrinkled quickly two or three times, so testing the frontalis muscle innervated by the facial nerve.
Gynaecomastia

Reference

Cross Reference
Facial paresis, Facial weakness

Gustatory Epiphora
- see CROCODILE TEARS

Gustatory Sweating
Gustatory sweating, Frey’s syndrome, is perspiration of the cheek, jaw, temple, and neck when eating, following aberrant regeneration of damaged autonomic fibres travelling in the glossopharyngeal and vagus nerves, for example, following neck dissection. This is an autonomic synkinesis. It may be treated with botulinum toxin injections.

Cross Reference
Synkinesis, Synkinesis

Guttmann’s Sign
Guttmann’s sign is facial vasodilatation associated with nasal congestion, hypertension, bradycardia, sweating, mydriasis, and piloerection, due to autonomic overactivity occurring as a feature of the acute phase of high spinal cord lesions.

Gynaecomastia
Gynaecomastia is inappropriate breast development in males. It may be observed in chronic liver disease and in certain neurological diseases:

- Excessive pituitary prolactin release secondary to impaired dopamine release from the hypothalamus due to local tumour or treatment with dopaminergic antagonist drugs (e.g. antipsychotic medications);
- Spinal and bulbar muscular atrophy (Kennedy’s syndrome, X-linked);
- Klinefelter’s syndrome;
- POEMS syndrome.
Habit Spasm
- see SPASM; TIC

Hallpike Manoeuvre, Hallpike Test
The Hallpike manoeuvre (Nylen–Bárány manoeuvre, positioning manoeuvre, Dix–Hallpike positioning test) is a test used in the investigation of vertigo to induce (or to modify) nystagmus by stimulating the otolith organs of the inner ear. It most usually consists of briskly tilting the patient’s head backwards to 30–45° below the horizontal (‘head hanging position’) and turning it 45° to one side or the other, thus stimulating the posterior semicircular canal. Prior to performing the manoeuvre, the examiner should warn the patient that s/he may feel ‘giddy’ or vertiginous, and to keep their eyes open throughout, since the development of nystagmus with the symptoms of vertigo is the observation of interest to the examiner.

With a peripheral lesion (e.g. benign paroxysmal positional vertigo, diseases of the labyrinth), nausea, vomiting, and rotational–vertical nystagmus occur several seconds after the manoeuvre and then rapidly fatigue (usually <30 s), only to recur when the patient is returned to the upright position, with the nystagmus now in the opposite direction. Repetition of the manoeuvre (if the patient can be persuaded to undergo it) causes less severe symptoms (habituation). This is the diagnostic test for benign paroxysmal positional vertigo (BPPV). Central lesions (disorders of the vestibular connections) tend to produce isolated nystagmus which does not fatigue or habituate with repetition.

Variants of the Hallpike manoeuvre are described for BPPV of anterior or horizontal semicircular canal origin. Caloric testing may be required to elicit the causes of dizziness if the Hallpike manoeuvre is uninformative.

References

Cross References
Caloric testing; Nystagmus; Vertigo; Vestibulo-ocular reflexes

Hallucination
A hallucination is a perception in the absence of adequate peripheral stimulus (cf. illusion, although there may be some overlap). Such perceptions are
Hallucination

substantial, constant, occur in objective space, and are usually not accompanied by insight (cf. pseudohallucination). They most usually occur in the visual and auditory domains.

Visual hallucinations may range in complexity. They may be ‘simple’, spots or flashes of light (photopsia, photism, scintillation), or ‘complex’, ranging from patterns (fortification spectra, epileptic aura) to fully formed objects or individuals. They may be transient, such as brief visions of a person or animal (passage hallucinations, for example, in Parkinson’s disease) or long lasting. Visual hallucinations may be normal, especially when falling asleep or waking (hypnagogic, hypnopompic). There are many other associations including both psychiatric and neurological disease, including

- Delirium: especially hyperalert/agitated subtype
- Withdrawal states: e.g. delirium tremens; hypnotics, anxiolytics
- Drug overdose: e.g. anticholinergic drugs
- Neurodegenerative disorders: dementia with Lewy bodies (a diagnostic criterion) more often than Alzheimer’s disease: these may be associated with cholinergic depletion and improved with cholinesterase inhibitor drugs; idiopathic Parkinson’s disease (with or without treatment)
- Narcolepsy–cataplexy
- Peduncular hallucinosis
- Migraine aura: usually visual or somaesthetic; less often auditory of olfactory
- Charles Bonnet syndrome (visual hallucinations of the visually impaired)
- Schizophrenia
- Epilepsy: complex partial seizures
- ‘Alice in Wonderland’ syndrome

Different mechanisms may account for visual hallucinations in different conditions: defective visual input and processing may occur in visual pathway lesions, whereas epilepsy may have a direct irritative effect on brain function; visual hallucinations associated with brainstem lesions may result from neurotransmitter abnormalities (cholinergic, serotonergic).

Auditory hallucinations may be simple (tinnitus) or complex (voices, music) and may be associated with focal pathology in the temporal cortex. Third person hallucinations, commenting on a person’s actions, are one of the first rank symptoms of schizophrenia.

References
Harvey’s Sign

A high-frequency tuning fork (440–1,024 Hz) applied to the mucosa overlying the nasal septum produces a painful stimulus which has been advocated as useful in differentiating true from simulated coma or status epilepticus.

Hammer Toes

Hammer toes are a feature of hereditary neuropathies, e.g. Charcot–Marie–Tooth disease, some cases of hereditary neuropathy with liability to pressure palsies, and Friedreich’s ataxia. There may be associated pes cavus.

Hand Elevation Test

This is one of the provocative tests for carpal tunnel syndrome: it is positive if paraesthesia in the distribution of the median nerve develop after raising the hand over the head for up to 2 min.

Harlequin Sign, Harlequin Syndrome

The harlequin sign or syndrome refers to asymmetrical facial flushing with sweating after exercise. That it reflects localized autonomic dysfunction may be indicated by its associations with congenital Horner’s syndrome, and as one element in the spectrum of Holmes–Adie syndrome and Ross’s syndrome. Harlequin sign has on occasion been described in association with multiple sclerosis and superior mediastinal neurinoma.

Harvey’s Sign

A high-frequency tuning fork (440–1,024 Hz) applied to the mucosa overlying the nasal septum produces a painful stimulus which has been advocated as useful in differentiating true from simulated coma or status epilepticus.
Hawthorne Effect

Hawthorne’s original observation was that children with learning disability only began to make progress when notice was taken of them, even though the intervention was unskilled or non-specific (which prompted Critchley to dare to wonder if this were the effect of speech therapy in chronic aphasics). The term Hawthorne effect has come to stand for any situation in which behaviour is altered by observation, or being the object of attention. In the neurological examination, certain signs may be evident when the patient is being observed, but absent when observation is only surreptitious, an inconsistency which may point to signs being ‘non-organic’, functional, or part of illness behaviour. A similar effect may be apparent in unblinded clinical trials.

References

Cross Reference
Functional weakness and sensory disturbance

Head Droop, Head Drop
- see DROPPED HEAD SYNDROME

Head Impulse Test
The head impulse test, also known as the head thrust test, assesses the vestibulo-ocular reflex. It consists of a rapid turning of the head to one side by about 15°, sufficiently rapid to ensure that smooth pursuit eye movements do not compensate for head turning. The examiner observes the ability of the subject to maintain fixation on a distant target; if the vestibulo-ocular reflex is intact fixation is maintained. If the vestibulo-ocular reflex is impaired, then an easily visible saccade back to the target occurs at the end of the movement. Tilting the head down by 20° and moving the head unpredictably may optimize testing. This test is recommended in patients suffering a first attack of acute spontaneous vertigo. Sensitivity and specificity of around 80% for detecting a peripheral vestibular lesion such as acute unilateral vestibular neuritis has been reported. To avoid false negatives, it has been suggested that the test should be performed with high acceleration, 5–10 times. If the test is normal in suspected vestibular neuritis, then a central cause such as cerebellar infarction needs to be excluded.

References


**Cross References**
Vertigo; Vestibulo-ocular reflexes

### Head Thrust
- see **EYELID APRAXIA; OCULAR APRAXIA**

### Head Tilt

Head tilt may be observed with:

- Diplopia, cranial nerve palsies (IV, VI); skew deviation
- Neck dystonia (laterocollis)
- Incipient tonsillar herniation with cerebellar tumours, sometimes associated
  with neck stiffness and limitation of neck movement

**Cross References**
Bielschowsky’s sign, Bielschowsky’s test; Diplopia; Laterocollis; Ocular tilt reaction

### Head Tremor

Head tremor may be characterized as ‘yes–yes’ (nodding, *tremblement affirmatif*) when predominantly in the vertical plane, or ‘no–no’ (side-to-side, *tremblement negatif*) when predominantly in the horizontal plane. Head tremor may occur in isolation or with evidence of tremor elsewhere (e.g. postural limb tremor, vocal tremor, in essential tremor), or dystonia (e.g. torticollis). In essential tremor the head movements are often intermittent, ‘yes–yes’, and of frequency about 7 Hz. Dystonic head tremor is often jerky and disorganized, with a frequency of less than 5 Hz. Cerebellum and brainstem disease such as multiple sclerosis can also produce head tremor (or titubation). Head tremor is an exceptionally rare symptom of Parkinson’s disease. It may also be seen as a consequence of aortic valve regurgitation (De Musset’s sign).

Treatment of head tremor varies with cause. Possible treatments, of variable efficacy, include

- **Essential tremor**: propranolol, topiramate, primidone, nicardipine, gabapentin;
- **Dystonic tremor**: levodopa, anticholinergics, propranolol, botulinum toxin injections;
- **Cerebellar tremor**: isoniazid, carbamazepine, ondansetron.

**Cross References**
Dystonia; Tremor

### Head Turning Sign

It is often observed that patients who are cognitively impaired turn their head towards their spouse, partner, or carer to seek assistance when asked to give a
Heautoscopy

This term was coined to denote seeing oneself, encountering one's alter ego or doppelgänger. Hence unlike the situation in autoscopy, there are two selves, a reduplicated body rather than a mirror image; egocentric and body-centred perspectives do not coincide. According to Critchley, the condition used to be called 'specular hallucinosis', and the Swedish naturalist Linnaeus (1707–1788) apparently had episodes, seeing himself sitting in his study or gazing at a flower and plucking it.

References


Cross References

Autoscopy; Hallucination

Heel–Knee–Shin Test, Heel–Shin Test

A frequently used test of coordination in which the patient, sitting on the examination couch, is asked to lift the heel onto the contralateral knee, then run it smoothly down the shin bone towards the foot. Jerky performance, or a tendency for the heel to slide off the shin, may be seen in an ataxic limb.

Cross References

Ataxia; Cerebellar syndromes; Shin-tapping

Heel–Toe Walking

- see TANDEM WALKING

Hemeralopia

Hemeralopia, or day blindness, is worsening of vision in bright light (cf. nyctalopia). This phenomenon may reflect severe impairment of blood flow to the eye, such that photostressing the macula by exposure to bright light is followed by only slow regeneration of the bleached photopigments. If due to retinal ischaemia, hemeralopia may be accompanied by neovascularization of the retina. Impoverished perfusion pressure may be demonstrated by pressing on the eyeball (e.g. with the thumb) during opthalmoscopy ('digital ophthalmodynamometry') and observing the collapse of retinal arteries: thumb pressure greater than diastolic retinal artery pressure causes intermittent collapse; thumb pressure greater than systolic pressure leads to a cessation of pulsation.

Hemeralopia may also occur in retinal diseases such as cone–rod dystrophies, and with cataract.
Hemianopia

Reference

Cross Reference
Nyctalopia

Hemiachromatopsia
- see ACHROMATOPSIA; ALEXIA

Hemiakinesia
Hemiakinesia is akinesia or hypokinesia (inability or difficulty initiating movement) confined to one side of the body. Although hemiakinesia is the norm at the onset of idiopathic Parkinson’s disease (‘hemiparkinsonism’), persistent hemiakinesia should prompt a re-evaluation of this diagnosis. Corticobasal degeneration often remains unilateral; a search for structural lesions of the basal ganglia should also be undertaken. Hemiakinesia may also indicate motor neglect, usually with right-sided lesions. Lesions of the basal ganglia, ventral (‘motor’) thalamus, limbic system, and frontal lobes may cause hemiakinesia.

Cross References
Akinesia; Extinction; Hemiparkinsonism; Hypokinesia; Neglect; Parkinsonism

Hemialexia
This is the inability to read words in the visual left half-field in the absence of hemianopia. It may occur after callosotomy (complete or partial involving only the splenium) and represents a visual disconnection syndrome.

Reference

Cross References
Alexia; Hemialexia

Hemianomia
This is the absence of verbal report of stimuli presented in the visual left half-field in the absence of hemianopia. It may occur after callosotomy (complete or partial involving only the splenium) and represents a visual disconnection syndrome.

Reference

Cross References
Anomia; Hemialexia

Hemianopia
Hemianopia (hemianopsia) is a defect of one-half of the visual field: this may be vertical or horizontal (= altitudinal field defect). Hemianopic defects may be congruent (homonymous) or non-congruent (heteronymous) and may be detected by
standard confrontational testing of the visual fields or by automated means (e.g. Goldman perimetry). These tests of the visual fields are an extension of the tests for visual acuity which assess areas away from the fovea. Because of the strict topographic arrangement of neural pathways within the visual system, particular abnormalities of the visual fields give a very precise indication of the likely site of pathology.

- **Homonymous hemianopia:**
  Reflects a postchiasmal lesion. It is important to assess whether the vertical meridian of a homonymous hemianopia cuts through the macula (macula splitting), implying a lesion of the optic radiation; or spares the macula (macula sparing), suggesting an occipital cortical lesion. Incongruous defects may be found with lesions of the optic tract. Commonly, homonymous hemianopias result from cerebrovascular disease causing occipital lobe infarction, or intraparanechymal tumour, but they may be ‘false-localizing’ due to raised intracranial pressure if temporal lobe herniation causes posterior cerebral artery compromise.

- **Heteronymous hemianopia:**
  Reflects a chiasmal lesion. The most common of these is a bitemporal hemianopia due to chiasmal compression, for example, by a pituitary lesion or craniopharyngioma. Tilted optic discs may also be associated with bitemporal field loss but this extends to the blind spot and not the vertical meridian as in chiasmal pathology (‘pseudobitemporal hemianopia’). Binasal defects are rare, suggesting lateral compression of the chiasm, for example, from bilateral carotid artery aneurysms; binasal hemianopia is also described with optic nerve head lesions. Unilateral (monocular) temporal hemianopia may result from a lesion anterior to the chiasm which selectively affects only the ipsilateral crossing nasal fibres (junctional scotoma of Traquair).

  Unawareness of visual field loss, anosognosic hemianopia, occurs principally with right-sided brain lesions.

  Bilateral homonymous hemianopia or double hemianopia may result in cortical blindness.

**Reference**

**Cross References**
Alexia; Altitudinal field defect; Anosognosia; Binasal hemianopia; Bitemporal hemianopia; Cortical blindness; ‘False-localizing signs’; Macula sparing, Macula splitting; Quadrantanopia; Scotoma; Visual field defects

**Hemiataxia**
Hemiataxia is ataxia confined to one-half of the body. The vast majority of isolated hemiataxic syndromes reflect a lesion of the ipsilateral cerebellar hemisphere, but on occasion supratentorial lesions may cause hemiataxia (posterior limb of the internal capsule, thalamus). However, in almost all of these cases
Hemiballismus

Hemiballismus is unilateral ballismus, an involuntary hyperkinetic movement disorder in which there are large amplitude, vigorous (‘flinging’) irregular movements. Hemiballismus overlaps clinically with hemichorea (‘violent chorea’); the term hemiballismus–hemichorea is sometimes used to reflect this overlap. Hemiballistic limbs may show a loss of normal muscular tone (hypotonia).

Neuroanatomically, hemiballismus is most often associated with lesions of the contralateral subthalamic nucleus of Luys or its efferent pathways, although there are occasional reports of its occurrence with lesions of the caudate nucleus, putamen, globus pallidus, lentiform nucleus, thalamus, and precentral gyrus; and even with ipsilateral lesions. Neuropathologically, vascular events (ischaemia, haemorrhage) are the most common association but hemiballismus has also been reported with space-occupying lesions (tumour, arteriovenous malformation), inflammation (encephalitis, systemic lupus erythematosus, post-streptococcal infection), demyelination, metabolic causes (hyperosmolal non-ketotic hyperglycaemia), infection (toxoplasmosis in AIDS), drugs (oral contraceptives, phenytoin, levodopa, neuroleptics), and head trauma.

Pathophysiologically, hemiballismus is thought to result from reduced conduction through the direct pathway within the basal ganglia–thalamo–cortical motor circuit (as are other hyperkinetic involuntary movements, such as choreothetosis). Removal of excitation from the globus pallidus following damage to the efferent subthalamic–pallidal pathways disinhibits the ventral anterior and ventral lateral thalamic nuclei which receive pallidal projections and which in turn project to the motor cortex.

Hemiballismus of vascular origin usually improves spontaneously, but drug treatment with neuroleptics (haloperidol, pimozide, sulpiride) may be helpful. Other drugs which are sometimes helpful include tetrabenazine, reserpine, clonazepam, clozapine, and sodium valproate.

References
Lee MS, Marsden CD. Movement disorders following lesions of the thalamus or subthalamic region. Movement Disorders 1994; 9: 493–507.
Martin JP. Hemichorea resulting from a local lesion of the brain. (The syndrome of the body of Luys.) Brain 1927; 50: 637–651.
Hemichorea
Hemichorea is unilateral chorea, an involuntary movement disorder which overlaps with hemiballismus, and with which it shares a similar pathophysiology and aetiology. It may replace hemiballismus during recovery from a contralateral subthalamic lesion.

Cross References
Chorea, Choreoathetosis; Hemiballismus

Hemidystonia
Hemidystonia is dystonia affecting the whole of one side of the body, a pattern which mandates structural brain imaging because of the chance of finding a causative structural lesion (vascular, neoplastic), which is greater than with other patterns of dystonia (focal, segmental, multifocal, generalized). Such a lesion most often affects the contralateral putamen or its afferent or efferent connections.

Reference

Cross Reference
Dystonia

Hemifacial Atrophy
- see PARRY–ROMBERG SYNDROME

Hemifacial Spasm
Hemifacial spasm is an involuntary dyskinetic (not dystonic) movement disorder consisting of painless contractions of muscles on one side of the face, sometimes triggered by eating or speaking, and exacerbated by fatigue or emotion. The movements give a twitching appearance to the eye or side of the mouth, sometimes described as a pulling sensation. Patients often find this embarrassing because it attracts the attention of others. The movements may continue during sleep. Paradoxical elevation of the eyebrow as orbicularis oris contracts and the eye closes may be seen (Babinski’s ‘other sign’). Very rarely, movements may be bilateral.

Hemifacial spasm may be idiopathic, or associated with neurovascular compression of the facial (VII) nerve, usually at the root entry zone, often by a tortuous anterior or posterior inferior cerebellar artery. Other causes include intrapontine lesions (e.g. demyelination), following a Bell’s palsy, and mass lesions (tumour, arteriovenous malformation) located anywhere from the facial nucleus to the stylomastoid foramen. Very rarely, contralateral (false-localizing) posterior fossa lesions have been associated with hemifacial spasm, suggesting that kinking or distortion of the nerve, rather than direct compression, may be of pathogenetic importance.
Hemiparesis

Structural lesions may be amenable to surgical resection. For idiopathic hemifacial spasm, or patients declining surgery, botulinum toxin injections are the treatment of choice.

Reference

Cross References
Babinski’s sign (2); Bell’s palsy; Dyskinesia; ‘False-localizing signs’

Hemi-Inattention
- see NEGLECT

Hemimicropsia
- see MICROPSIA

Hemineglect
- see NEGLECT

Hemiparesis
Hemiparesis is a weakness affecting one side of the body, less severe than a hemiplegia. Characteristically this affects the extensor muscles of the upper limb more than flexors, and the flexors of the leg more than extensors (‘pyramidal’ distribution of weakness), producing the classic hemiparetic/hemiplegic posture with flexed arm and extended leg, the latter permitting standing and a circumducting gait.

Hemiparesis results from damage (most usually vascular) to the corticospinal pathways anywhere from motor cortex to the cervical spine. Accompanying signs may give clues as to localization, the main possibilities being hemisphere, brainstem, or cervical cord.

Hemisphere lesions may also cause hemisensory impairment, hemianopia, aphasia, agnosia, or apraxia; headache, and incomplete unilateral ptosis, may sometimes feature. Spatial neglect, with or without anosognosia, may also occur, particularly with right-sided lesions producing a left hemiparesis. Pure motor hemiparesis may be seen with lesions of the internal capsule, corona radiata, and basal pons (lacunar/small deep infarct), in which case the face and arm are affected more than the leg; such facio-brachial predominance may also be seen with cortico–subcortical lesions laterally placed on the contralateral hemisphere. Crural predominance suggests a contralateral paracentral cortical lesion or one of the lacunar syndromes.

Brainstem lesions may produce diplopia, ophthalmoplegia, nystagmus, ataxia, and crossed facial sensory loss or weakness in addition to hemiparesis (‘alternating hemiplegia’).

Spinal lesions are more likely to show bilateral long tract signs (e.g. bilateral Babinski’s sign) and may have accompanying spinal or root pain, sphincter disturbance, and a sensory or motor level.

Hemiparesis is most usually a consequence of a vascular event (cerebral infarction). Tumour may cause a progressive hemiparesis (although meningiomas may produce transient ‘stroke-like’ events). Hemiparetic multiple sclerosis is rare but well described. Transient hemiparesis may be observed as
Hemiparkinsonism

Hemiparkinsonism describes the finding of parkinsonian signs restricted to one side of the body, most usually akinesia, in which case the term hemiakinesia may be used. Idiopathic Parkinson’s disease may present with exclusively or predominantly unilateral features (indeed, lack of asymmetry at onset may argue against this diagnosis) but persistent hemiparkinsonism, particularly if unresponsive to adequate doses of levodopa, should alert the clinician to other possible diagnoses, including corticobasal degeneration or structural lesions.

Cross References
Hemiakinesia; Parkinsonism

Hemiplegia

Hemiplegia is a complete weakness affecting one side of the body, i.e. clinically a more severe picture than hemiparesis.

Cross References
Hemiparesis; Weakness

Hemiplegia Cruciata

Cervico-medullary junction lesions where the pyramidal tract decussates may result in paresis of the contralateral upper extremity and ipsilateral lower extremity. There may be concurrent facial sensory loss with onion skin pattern, respiratory insufficiency, bladder dysfunction, and cranial nerve palsies. Such cases are very rare.

Cross Reference
Onion peel, Onion skin

Hennebert’s Sign

Hennebert’s sign is the induction of vertigo and nystagmus by pressure changes in the external auditory canal, such as when using pneumatic otoscopy or simply with tragal pressure. These findings are highly suggestive of the presence of a bony labyrinthine fistula. There may be a history of chronic otitis media.

Cross References
Nystagmus; Vertigo

Henry and Woodruff Sign

Evidence of visual fixation, reported to be helpful in differentiating pseudo-seizures from epileptic seizures: the patient is rolled from one side on to the other whilst note is taken of whether the eyes remain directed towards the ground.
Heterotropia

References

Hertwig–Magendie Sign
- see SKEW DEVIATION

Heterochromia Iridis
Different colour of the irides may be seen in congenital Horner’s syndrome and in Waardenburg syndrome of nerve deafness, white forelock, abnormal skin pigmentation, and synophrys.

Cross Reference
Horner’s syndrome

Heterophoria
Heterophoria is a generic term for a latent tendency to imbalance of the ocular axes (latent strabismus; cf. heterotropia). This may be clinically demonstrated using the cover–uncover test: if there is movement of the covered eye as it is uncovered and takes up fixation, this reflects a phoria. Phorias may be in the horizontal (esophoria, exophoria) or vertical plane (hyperphoria, hypophoria).

Reference

Cross References
Cover tests; Esophoria; Exophoria; Heterotropia; Hyperphoria; Hypophoria

Heterotropia
Heterotropia is a generic term for manifest deviation of the eyes (manifest strabismus; cf. heterophoria), synonymous with squint. This may be obvious; an amblyopic eye, with poor visual acuity and fixation, may become deviated. Sometimes it may be more subtle, coming to attention only with the patient’s complaint of diplopia.

Using the alternate cover (cross-cover) test, in which binocular fixation is not permitted, an imbalance in the visual axes may be demonstrated, but this will not distinguish between heterotropia and heterophoria. To make this distinction the cover test is required: if the uncovered eye moves to adopt fixation then heterotropia is confirmed. Tropias may be in the horizontal (esotropia, exotropia) or vertical plane (hypertropia, hypotropia).

Reference

Cross References
Amblyopia; Cover tests; Esotropia; Exotropia; Heterophoria; Hypertropia; Hypotropia

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Hiccups

A hiccup (hiccough) is a brief burst of inspiratory activity involving the diaphragm and the inspiratory intercostal muscles with reciprocal inhibition of expiratory intercostal muscles. The sound (‘hic’) and discomfort result from glottic closure immediately after the onset of diaphragmatic contraction, i.e. the latter is insufficient or asynchronous. Hiccups may be characterized as a physiological form of myoclonus (or singultus).

Most episodes of hiccups are self-limited, but prolonged or intractable hiccuping (hocquet diabolique) should prompt a search for a structural or functional cause, either gastroenterological or neurological. Hiccuping is seldom the only abnormality if the cause is neurological since it usually reflects pathology within the medulla or affecting the afferent and efferent nerves of the respiratory muscles. Medullary causes include

- Infarction (posterior inferior cerebellar artery territory; lateral medullary syndrome, especially middle level and dorsolateral lesion locations)
- Tumour
- Abscess
- Tuberculoma
- Syrinx
- Haematoma
- Demyelination
- CNS infection, e.g. viral encephalitis

Treatment should be aimed at the underlying cause. If none is identified, physical measures to stop the hiccups such as rebreathing may then be tried. Of the many various pharmacotherapies tried, the best are probably baclofen and chlorpromazine.

References


Cross References

Lateral medullary syndrome; Myoclonus

High-Stepping Gait

- see STEPPAGE, STEPPING GAIT

Hip Abduction Sign

The hip abduction sign refers to abduction of the thighs when attempting to rise from the ground, due to relative weakness of hip adductors with preserved strength in hip abductors. The sign was first described in patients with sarcoglycanopathies, a group of autosomal recessive limb-girdle muscular dystrophies,
and is reported to have a sensitivity of 76% and a specificity of 98% for this diagnosis. It may perhaps be envisaged as the equivalent to Gowers’ sign but with hip adductor, rather than gluteal, weakness.

Reference

Cross Reference
Gowers’ sign

Hippus
Hippus is excessive pupillary unrest, i.e. rhythmic, oscillatory, contraction and dilatation of the pupil. It may reflect an imbalance between afferent pupillary sympathetic and parasympathetic autonomic activity. Hippus may be a normal phenomenon; it may be observed during recovery from an oculomotor (III) nerve palsy, but otherwise is of no localizing significance.

Hitselberg Sign
Hypoesthesia of the posterior wall of the external auditory canal may be seen in facial paresis since the facial nerve sends a sensory branch to innervate this territory.

Cross Reference
Facial paresis, Facial weakness

Hocquet Diabolique
- see HICCUPS

Hoffmann’s Sign
Hoffmann’s sign or reflex is a digital reflex consisting of flexion of the thumb and index finger in response to snapping or flicking the distal phalanx of the middle finger, causing a sudden extension of the joint. Although sometimes a normal finding, for example, in the presence of generalized hyperreflexia (anxiety, hyperthyroidism), it may be indicative of a corticospinal tract lesion above C5 or C6, particularly if present unilaterally.

Cross References
Trömner’s sign; Upper motor neurone (UMN) syndrome

Hoffmann–Tinel Sign
- see TINEL’S SIGN

Holmes–Adie Pupil, Holmes–Adie Syndrome
The Holmes–Adie, or tonic, pupil is an enlarged pupil which, in a darkened environment, is unresponsive to a phasic light stimulus, but may respond slowly to a tonic light stimulus. Reaction to accommodation is preserved (partial iridoplegia), hence this is one of the causes of light-near pupillary dissociation. A Holmes–Adie pupil is usually unilateral, and hence a cause of anisocoria.

A Holmes–Adie pupil may be associated with other neurological features (Holmes–Adie syndrome). These include loss of lower limb tendon reflexes (especially ankle jerks); impaired corneal sensation; chronic cough; and localized
or generalized anhidrosis, sometimes with hyperhidrosis (Ross’s syndrome). Holmes–Adie syndrome is much more common in women than men.

Pathophysiologically Holmes–Adie pupil results from a peripheral lesion of the parasympathetic autonomic nervous system and shows denervation supersensitivity, constricting with application of dilute (0.2%) pilocarpine (cf. pseudo-Argyll Robertson pupil).

References

Cross References
Anhidrosis; Anisocoria; Hyperhidrosis; Light-near pupillary dissociation; Pseudo-Argyll Robertson pupil

Holmes’ Tremor
Holmes’ tremor, also known as rubral tremor, or midbrain tremor, has been defined as a rest and intention tremor, of frequency <4.5 Hz. The rest tremor may resemble parkinsonian tremor and is exacerbated by sustained postures and voluntary movements. Hence there are features of rest, postural and kinetic (intention) tremor. Once attributed to lesions of the red nucleus (hence ‘rubral’), the anatomical substrate is now thought to be interruption of fibres of the superior cerebellar peduncle (hence ‘midbrain’) carrying cerebellothalamic and/or cerebello-olivary projections; lesions of the ipsilateral cerebellar dentate nucleus may produce a similar clinical picture. Recognized causes include multiple sclerosis, head injury, and stroke. If a causative lesion is defined, there is typically a delay before tremor appearance (4 weeks to 2 years).

References

Cross Reference
Tremor

Hoover’s Sign
Hoover’s sign may be used to help differentiate organic from functional hemiplegia or monoplegia. It is based on the fact that when a recumbent patient attempts to lift one leg, downward pressure is felt under the heel of the other leg, hip extension being a normal synergistic or synkinesis movement. The finding of this synkinetic movement, detected when the heel of the supposedly paralyzed leg presses down on the examiner’s palm, constitutes Hoover’s sign: no increase in pressure is felt beneath the heel of a paralyzed leg in an organic hemiplegia.
Horner’s Syndrome

In addition, the synkinetic hip extension movement is accentuated when attempting to raise a contralateral paretic leg, whereas in functional weakness it is abolished.

Reference

Cross References
‘Arm drop’; Babinski’s trunk–thigh test; Functional weakness and sensory disturbance; Synkinesia, Synkinesis

Horner’s Syndrome

Horner’s syndrome, or Bernard–Horner syndrome, is defined by a constellation of clinical findings, most usually occurring unilaterally, viz.:

- partial ptosis, due to weakness of Müller’s muscle;
- miosis, due to the unopposed action of the sphincter pupillae muscle, innervated by the parasympathetic nervous system; this is most obvious in a dimly lit room;
- anhidrosis, a loss of sweating (if the lesion is distal to the superior cervical ganglion);
- enophthalmos, retraction of the eyeball (though this is seldom measured).

The first two mentioned signs are usually the most evident and bring the patient to medical attention; the latter two are usually less evident or absent. Additional features which may be seen include

- heterochromia iridis, different colour of the iris (if the lesion is congenital);
- elevation of the inferior eyelid due to a weak inferior tarsal muscle (‘reverse ptosis’ or ‘upside-down ptosis’).

Horner’s syndrome results from impairment of ocular sympathetic innervation. The sympathetic innervation of the eye consists of a long, three neurone, pathway, extending from the diencephalon down to the cervicothoracic spinal cord, then back up to the eye via the superior cervical ganglion and the internal carotid artery, and the ophthalmic division of the trigeminal (V) nerve. A wide variety of pathological processes, spread across a large area, may cause a Horner’s syndrome, although many examples remain idiopathic despite intensive investigation. Hence Horner’s syndrome is a good lateralizing but a poor localizing sign. Recognized causes include

- brainstem/cervical cord disease (vascular, demyelination, syringomyelia);
- Pancoast tumour;
- malignant cervical lymph nodes;
- carotid aneurysm, carotid artery dissection;
- involvement of T1 fibres, e.g. in T1 radiculopathy, or lower trunk brachial plexopathy;
- cluster headache;
- congenital.

Determining whether the lesion causing a Horner’s syndrome is preganglionic or postganglionic may be done by applying to the eye 1% hydroxyamphetamine hydrobromide, which releases noradrenaline into the synaptic cleft, which dilates the pupil if Horner’s syndrome results from a preganglionic lesion.
However, this is not particularly helpful in determining cause, whereas accompanying neurological features are as follows: contralateral hemiparesis would mandate investigation for carotid dissection (MRI, MRA, angiography), and this is probably sensible for any painful Horner’s syndrome of acute onset. Arm symptoms and signs in a smoker mandate a chest radiograph for Pancoast tumour. If the Horner’s syndrome is isolated and painless, then no investigation may be required. In this situation, a symptomatic cause is seldom identified despite investigation. Syringomyelia presenting with isolated Horner’s syndrome has been reported.

Unilateral miosis may be mistaken for contralateral mydriasis if ptosis is subtle, leading to suspicion of a partial oculomotor nerve palsy on the ‘mydriatic’ side. Observation of anisocoria in the dark will help here, since increased anisocoria indicates a sympathetic defect (normal pupil dilates) whereas less anisocoria suggests a parasympathetic lesion. Applying to the eye 10% cocaine solution will also diagnose a Horner’s syndrome if the pupil fails to dilate after 45 min in the dark (normal pupil dilates).

Reference

Cross References
Anhidrosis; Anisocoria; Enophthalmos; Miosis; Plexopathy; Ptosis; Radiculopathy

**Hoyt–Spencer Sign**
This name is given to the triad of findings characteristic of chronic optic nerve compression, especially due to sphenoorbital optic nerve sheath meningiomas:

- Optociliary shunt vessels
- Disc pallor
- Visual loss

‘Hung-Up’ Reflexes
- see WOLTMAN’S SIGN

**Hutchinson’s Pupil**
Hutchinson’s pupil is unilateral pupillary dilatation ipsilateral to a supratentorial (usually extrinsic) space-occupying lesion, which may be the earliest sign of raised intracranial pressure. It reflects involvement of peripheral pupilloconstrictor fibres in the oculomotor (III) nerve, perhaps due to compression on the margin of the tentorium.

**Cross References**
Anisocoria; Mydriasis; Oculomotor (III) nerve palsy

**Hyperacusis**
Hyperacusis is an abnormal loudness of sounds, especially low tones, due to paralysis of the stapedius muscle, whose normal reflex function is to damp conduction across the ossicular chain of the middle ear. This most commonly occurs with lower motor neurone facial (VII) nerve (Bell’s) palsy, located proximal to

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Hyperekplexia

the nerve to stapedius. Ageusia may also be present if the chorda tympani branch of the facial nerve is involved. Hyperacusis may occasionally occur with central (brainstem) lesions.

Reduction or absence of the stapedius reflex may be tested using the stethoscope loudness imbalance test: with a stethoscope placed in the patients ears, a vibrating tuning fork is placed on the bell. Normally the perception of sound is symmetrical, but sound lateralizes to the side of facial paresis if the attenuating effect of the stapedius reflex is lost.

Cross References
Ageusia; Bell’s palsy; Facial paresis, Facial weakness

Hyperaesthesia
Hyperaesthesia is increased sensitivity to sensory stimulation of any modality, e.g. pain (hyperalgesia), touch.

Cross References
Anaesthesia; Hyperalgesia

Hyperalgesia
Hyperalgesia is the exaggerated perception of pain from a stimulus which is normally painful (cf. allodynia). This may result from sensitization of nociceptors (paradoxically this may sometimes be induced by morphine) or abnormal ephaptic cross-excitation between primary afferent fibres.

Cross References
Allodynia; Dysaesthesia; Hyperpathia

Hyperekplexia
Hyperekplexia (literally, to jump excessively) is an involuntary movement disorder in which there is a pathologically exaggerated startle response, usually to sudden unexpected auditory stimuli, but sometimes also to tactile (especially trigeminal) and visual stimuli. The startle response is a sudden shock-like movement which consists of eye blink, grimace, abduction of the arms, and flexion of the neck, trunk, elbows, hips, and knees. The muscular jerk of startle satisfies the definition of myoclonus. Ideally for hyperekplexia to be diagnosed there should be a physiological demonstration of exaggerated startle response, but this criterion is seldom adequately fulfilled. Hyperekplexia syndromes may be classified as:

- **Idiopathic**: the majority.
- **Hereditary/familial**: An autosomal dominant disorder with muscular hypertonia in infancy, leg jerks, and gait disorder. Familial cases have been associated with mutations in the α1 subunit of the inhibitory glycine receptor gene.
- **Symptomatic**: perinatal ischaemic–hypoxic encephalopathy; brainstem lesions (encephalitis, haemorrhage); thalamic lesions (inflammation, vascular); drugs (cocaine, amphetamines); Tourette syndrome.
Hypergraphia

Attacks may respond to the GABA agonist clonazepam.

References

Cross References
Incontinence; Myoclonus

Hypergraphia

Hypergraphia is a form of increased writing activity. It has been suggested that it should refer specifically to all transient increased writing activity with a non-iterative appearance at the syntactic or lexicographemic level (cf. automatic writing behaviour).

Hypergraphia may be seen as part of the interictal psychosis which sometimes develops in patients with complex partial seizures from a temporal lobe (especially non-dominant hemisphere) focus, or with other non-dominant temporal lobe lesions (vascular, neoplastic, demyelinating, neurodegenerative), or psychiatric disorders (schizophrenia). Hypergraphia is a feature of Geschwind’s syndrome, along with hyperreligiosity and hyposexuality.

References

Cross References
Automatic writing behaviour; Hyperreligiosity; Hyposexuality

Hyperhidrosis

Hyperhidrosis is excessive (unphysiological) sweating. This may be ‘essential’ (i.e. without obvious cause), or seen as a feature of acromegaly, Parkinson’s disease, or occurring in a band above a spinal cord injury. Localized hyperhidrosis caused by food (gustatory sweating) may result from aberrant connections between nerve fibres supplying sweat glands and salivary glands. Other causes of hyperhidrosis include mercury poisoning, phaeochromocytoma, and tetanus. Transient hyperhidrosis contralateral to a large cerebral infarct in the absence of autonomic dysfunction has also been described. Regional syndromes of hyperhidrosis (hands, feet, axillae) are also described.

Treatment is difficult. Symptoms may be helped (but not abolished) by low dose anticholinergic drugs, clonidine, or propantheline. For focal syndromes, botulinum toxin injections or sympathectomy may be helpful.

References
Hyperpathia

Cross References
Anhidrosis; Diaphoresis; Gustatory sweating; Holmes–Adie pupil, Holmes–Adie syndrome

Hyperkinesia
Hyperkinesia indicates an involuntary movement disorder characterized by excessive amplitude of movement, such as ballism, or chorea, or the speech disorders occurring with them.

Cross References
Ballism, Ballismus; Chorea, Choreoathetosis; Dysarthria

Hyperlexia
Hyperlexia has been used to refer to the ability to read easily and fluently (i.e. decode printed text to sound) without necessarily comprehending the meaning of the text which has been read, not infrequent in cases of autism.

Hypermetamorphosis
Hypermetamorphosis is an over attention to external stimuli. Patients with hypermetamorphosis may explore compulsively and touch everything in their environment. This is one element of the environmental dependency syndrome and may be associated with other forms of utilization behaviour, imitation behaviour (echolalia, echopraxia), and frontal release signs such as the grasp reflex. It occurs with severe frontal lobe damage and may be observed following recovery from herpes simplex encephalitis and in frontal lobe dementias including Pick’s disease. Bitemporal lobectomy may also result in hypermetamorphosis, as a feature of the Klüver–Bucy syndrome.

Cross References
Attention; Echolalia; Echopraxia; Frontal release signs; Grasp reflex; Imitation behaviour; Klüver–Bucy syndrome; Utilization behaviour

Hypermetria
- see DYSMETRIA

Hypermnesia
- see EIDETIC MEMORY; SYNAESTHESIA

Hyperorality
Hyperorality is a neurobehavioural abnormality consisting of drinking more than usual, eating excessively, eating anything in sight, and putting objects inappropriately into the mouth. It is a feature of frontal lobe pathology. It is one element of the Klüver–Bucy syndrome, along with hypersexuality.

Cross References
Geophagia, Geophagy; Klüver–Bucy syndrome

Hyperpathia
Hyperpathia is an unpleasant sensation, often a burning pain, associated with elevated threshold for cutaneous sensory stimuli such as light touch or hot and cold stimuli, especially repetitive stimuli. Even light stimuli may produce pain. Clinical features of hyperpathia may include summation (pain perception
increases with repeated stimulation) and aftersensations (pain continues after stimulation has ceased). The term thus overlaps to some extent with hyperalgesia (although the initial stimulus need not be painful itself) and dysesthesia. There is an accompanying diminution of sensibility due to raising of the sensory threshold (cf. allodynia), and the pain is not stimulus-bound (i.e. spreads beyond the area of stimulation).

Hyperpathia is a feature of thalamic lesions, and hence tends to involve the whole of one side of the body following a unilateral lesion such as a cerebral haemorrhage or thrombosis. Generalized hyperpathia may also be seen in variant Creutzfeldt–Jakob disease, in which posterior thalamic (pulvinar) lesions are said to be a characteristic neuroradiological finding.

**Cross References**

Allodynia; Dysesthesia; Hyperalgesia

**Hyperphagia**

Hyperphagia is increased or excessive eating. Binge eating, particularly of sweet things, is one of the neurobehavioural disturbances seen in certain of the frontotemporal dementias. Hyperphagia may be one feature of a more general tendency to put things in the mouth (hyperorality), for example, in the Klüver–Bucy syndrome.

**Cross References**

Hyperorality; Klüver–Bucy syndrome

**Hyperphoria**

Hyperphoria is a variety of heterophoria in which there is a latent upward deviation of the visual axis of one eye. Using the cover–uncover test, this may be observed clinically as the downward movement of the eye as it is uncovered.

**Cross References**

Cover tests; Heterophoria; Hypophoria

**Hyperpilaphesie**

The name given to the augmentation of tactile faculties in response to other sensory deprivation, for example, touch sensation in the blind.

**Hyperpronation**

- see CHOREA, CHOREOATHETOSIS; DECEREBRATE RIGIDITY

**Hyperreflexia**

Hyperreflexia is an exaggerated briskness of the tendon reflexes. This may be physiological in an anxious patient (reflexes often denoted ++), or pathological in the context of corticospinal pathway pathology (upper motor neurone syndrome, often denoted +++). It is sometimes difficult to distinguish normally brisk reflexes from pathologically brisk reflexes. Hyperreflexia (including a jaw jerk) in isolation cannot be used to diagnose an upper motor neurone syndrome, and asymmetry of reflexes is a soft sign. On the other hand, upgoing plantar responses are a hard sign of upper motor neurone pathology; other accompanying signs (weakness, sustained clonus, and absent abdominal reflexes) also indicate abnormality.
Hypersomnolence

Hypersomnolence is characterized by excessive daytime sleepiness, with a tendency to fall asleep at inappropriate times and places, for example, during

Hyperreflexia reflects an increased gain in the stretch reflex. This may be due to impaired descending inhibitory inputs to the monosynaptic reflex arc. Rarely pathological hyperreflexia may occur in the absence of spasticity, suggesting different neuroanatomical substrates underlying these phenomena.

‘Hyperreflexia’ of the bladder detrusor muscle may be a cause of urinary urge incontinence

Reference

Cross References
Abdominal reflexes; Clonus; Incontinence; Jaw jerk; Reflexes; Spasticity; Upper motor neurone (UMN) syndrome; Weakness

Hyperreligiosity

Hyperreligiosity is a neurobehavioural symptom, manifest as sudden religious conversion, or increased and unswerving orthodoxy in devotion to religious rituals. It may be encountered along with hypergraphia and hyposexuality as a feature of Geschwind’s syndrome. It has also been observed in some patients with frontotemporal dementia; the finding is cross-cultural, having been described in Christians, Muslims, and Sikhs. In the context of refractory epilepsy, it has been associated with reduced volume of the right hippocampus, but not right amygdala.

References

Cross References
Hypergraphia; Hyposexuality

Hypersexuality

Hypersexuality is a pathological increase in sexual drive and activity. Recognized causes include bilateral temporal lobe damage, as in the Klüver–Bucy syndrome, septal damage, hypothalamic disease (rare) with or without subjective increase in libido, and dopaminergic drug treatment in Parkinson’s disease. Hypersexuality is also a feature of the Kleine–Levin syndrome. Sexual disinhibition may be a feature of frontal lobe syndromes, particularly of the orbitofrontal cortex.

Reference

Cross References
Disinhibition; Frontal lobe syndromes; Klüver–Bucy syndrome; Punding

Hypersonmnoleance

Hypersonmnoleance is characterized by excessive daytime sleepiness, with a tendency to fall asleep at inappropriate times and places, for example, during
meals, telephone conversations, at the wheel of a car. Causes of hypersomnolence include

- Narcolepsy or the narcoleptic syndrome: may be accompanied by other features such as sleep paralysis, hypnagogic hallucinations, cataplexy
- Midbrain lesions
- Idiopathic CNS hypersomnia
- Kleine–Levin syndrome
- Nocturnal hypoventilation, due to:
  - Obstructive sleep apnoea–hypopnoea syndrome (OSAHS; Pickwickian syndrome)
  - Chest wall anomalies
  - Neuromuscular and myopathic disorders affecting the respiratory muscles, especially the diaphragm, for example:
    - motor neurone disease
    - myotonic dystrophy
    - metabolic myopathies, e.g. acid maltase deficiency
    - mitochondrial disorders
- Drugs: benzodiazepines, ergot-derivative dopamine agonists
- Post-stroke sleep-related disorders

Nocturnal hypoventilation as a consequence of obstructed breathing, often manifest as snoring, causes arterial oxygen desaturation as a consequence of hypopnoea/apnoea which may lead to disturbed sleep, repeated arousals associated with tachycardia, and hypertension. Clinical signs may include a bounding hyperdynamic circulation and sometimes papilloedema, as well as features of any underlying neuromuscular disease. OSAHS may present in the neurology clinics with loss of consciousness (sleep secondary to hypersomnolence), stroke, morning headaches, and cognitive impairment (slowing). Investigations may reveal a raised haematocrit and early morning hypoxia. Sleep studies confirm nocturnal hypoventilation with dips in arterial oxygen saturation. Treatment is with nocturnal intermittent positive pressure ventilation. Modafinil is also licensed for this indication.

Cross References
Asterixis; Cataplexy; Papilloedema; Paradoxical breathing; Snoring

Hyperthermia

Body temperature is usually regulated within narrow limits through the coordinating actions of a centre for temperature control (‘thermostat’), located in the hypothalamus (anterior–preoptic area), and effector mechanisms (shivering, sweating, panting, vasoconstriction, vasodilation), controlled by pathways located in or running through the posterior hypothalamus and peripherally in the autonomic nervous system. Lesions of the anterior hypothalamus (e.g. trauma, ischaemia, inflammation, tumour) may result in hyperthermia (cf. hypothermia). Other recognized causes of hyperthermia include

- Infection: bacteria, viruses (pyrogens, e.g. interleukin-1)
- Malignant hyperthermia
Hypohidrosis

- Neuroleptic malignant syndrome (hyperthermia, rigidity, autonomic dysfunction)
- Heatstroke
- Hyperthyroidism
- Phaeochromocytoma crisis

Cross Reference
Hypothermia

Hypertonia, Hypertonus
Hypertonia or hypertonus is an exaggeration of normal muscular tone, manifest as resistance to passive movement. It usually implies spasticity of corticospinal (pyramidal) pathway origin, rather than (leadpipe) rigidity of extrapyramidal origin.

Cross References
Clasp-knife phenomenon; Hyperreflexia; Paratonia; Rigidity; Spasiticity; Upper motor neurone (UMN) syndrome

Hypertrophy
- see MUSCLE HYPERTROPHY

Hypertropia
Hypertropia is a variety of heterotropia in which there is manifest upward vertical deviation of the visual axis of one eye. Using the cover test, this manifests as downward movement of the uncovered eye. Depending on the affected eye, this finding is often described as a ‘left-over right’ or ‘right-over left’. Asymptomatic hypertropia on lateral gaze is often congenital or physiological.

Cross References
Bielschowsky’s sign, Bielschowsky’s test; Cover tests; Heterotropia; Hypotropia

Hypoesthesia
Hypoesthesia (hypesthesia) is increased sensitivity to, or diminution of, sensory perception in any modality, most frequently used to describe pain (hypalgesia) or touch.

Cross Reference
Anaesthesia

Hypalgesia
Hypalgesia is a decreased sensitivity to, or diminution of, pain perception in response to a normally painful stimulus.

Cross Reference
Analgesia

Hypoguesia
- see AGEUSIA

Hypohidrosis
- see ANHIDROSIS
Hypokinesia
Hypokinesia is a reduction in the speed of initiation of voluntary movements, which at worst may progress to an inability to initiate voluntary movement (akinesia). Repeated apposition of finger and thumb or foot tapping may be useful in demonstrating hypokinesia of gradual onset (‘fatigue’). It may often coexist with bradykinesia and hypometria and is a feature of disorders of the basal ganglia (akineti-rigid or parkinsonian syndromes), for example:

- Parkinson’s disease
- Multiple system atrophy
- Progressive supranuclear palsy (Steele–Richardson–Olszewski syndrome)
- Some variants of prion disease

Cross References
Akinesia; Bradykinesia; Fatigue; Parkinsonism

Hypometria
Hypometria is a reduction in the amplitude of voluntary movements. It may be demonstrated by asking a patient to make repeated, large amplitude, opposition movements of thumb and forefinger, or tapping movements of the foot on the floor. A gradual decline in amplitude (which may be referred to as fatiguability; cf. fatigue) denotes hypometria. Voluntary saccadic eye movements may also show a ‘step’, as a correcting additional saccade compensates for the undershoot (hypometria) of the original movement. Hypometria is a feature of parkinsonian syndromes such as idiopathic Parkinson’s disease.

Cross References
Akinesia; Bradykinesia; Dysmetria; Fatigue; Hypokinesia; Parkinsonism; Saccades

Hypomimia
Hypomimia, or amimia, is a deficit or absence of expression by gesture or mimicry. This is usually most obvious as a lack of facial expressive mobility (‘mask-like facies’). This is a feature of frontal–subcortical disease, e.g. basal ganglia disease producing akinetic-rigid or parkinsonian syndromes, and frontal lobe lesions (especially of the non-dominant hemisphere).

Cross References
Facial paresis, Facial weakness; Fisher’s sign; Parkinsonism

Hypophonia
Hypophonia is a quiet voice, as in hypokinetic dysarthria. It is often a feature of parkinsonian syndromes (e.g. idiopathic Parkinson’s disease, multiple system atrophy) and may occur early in progressive supranuclear palsy. In isolation, other causes of dysphonia may need to be considered.

Cross References
Dysarthria; Dysphonia; Parkinsonism

Hypophoria
Hypophoria is a variety of heterophoria in which there is a latent downward deviation of the visual axis of one eye. Using the cover–uncover test, this may be observed clinically as the upward movement of the eye as it is uncovered.

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Hypoflexia
Hypoflexia is a diminution of tendon reflexes, short of their total absence (areflexia). This may be physiological, as with the diminution of the ankle jerks with normal ageing; or pathological, most usually as a feature of peripheral lesions such as radiculopathy or neuropathy. The latter may be axonal or demyelinating, in the latter the blunting of the reflex may be out of proportion to associated weakness or sensory loss. Although frequently characterized as a feature of the lower motor neurone syndrome, the pathology underlying hypoflexia may occur anywhere along the monosynaptic reflex arc, including the sensory afferent fibre and dorsal root ganglion as well as the motor efferent fibre, and/or the spinal cord synapse.

Hypoflexia may also accompany central lesions, particularly with involvement of the mesencephalic and upper pontine reticular formation. Hypoflexia is an accompaniment of hemiballismus, and may also be noted in brainstem encephalitis (Bickerstaff’s encephalitis), in which the presence of a peripheral nerve disorder is debated. Hypoflexia is not a feature of myasthenia gravis but may occur in Lambert–Eaton myasthenic syndrome (cf. facilitation); it is not seen in most muscle diseases unless they are advanced.

Cross References
Age-related signs; Areflexia; Facilitation; Lower motor neurone (LMN) syndrome; Reflexes

Hyposexuality
Hyposexuality is a lack of sexual drive, interest, or activity. It may be associated with many diseases, physical or psychiatric, and/or medications which affect the central nervous system. Along with hypergraphia and hyperreligiosity, hyposexuality is one of the defining features of the Geschwind syndrome.

References

Cross References
Hypergraphia; Hyperreligiosity

Hypothermia
Hypothalamic damage, particularly in the posterior region, can lead to hypothermia (cf. hyperthermia) or poikilothermia (body temperature varying with ambient temperature, as in reptiles). There are many pathological causes, including tumour, trauma, infarct, haemorrhage, neurosarcoidosis, Wernicke’s encephalopathy, fat embolism, histiocytosis X, and multiple sclerosis (rare). A rare syndrome of paroxysmal or periodic hypothermia has been described and labelled as diencephalic epilepsy. Non-neurological causes of hypothermia are more common, including hypothyroidism, hypopituitarism, hypoglycaemia, and drug overdose.
Hypotonia, Hypotonus

Reference

Cross Reference
Hyperthermia

**Hypotonia, Hypotonus**
Hypotonia (hypotonus) is a diminution or loss of normal muscular tone, causing flappiness of the limbs. This is particularly associated with peripheral nerve or muscle pathology, as well as lesions of the cerebellum and certain basal ganglia disorders such as hemiballismus–hemichorea. Weakness preventing voluntary activity rather than a reduction in stretch reflex activity appears to be the mechanism of hypotonia.

Reference

Cross References
Ataxia; Flaccidity; Hemiballismus; Hypertonia

**Hypotropia**
Hypotropia is a variety of heterotropia in which there is manifest downward vertical deviation of the visual axis of one eye. Using the cover test, this manifests as upward movement of the uncovered eye. Depending on the affected eye, this finding is often described as a ‘left-over-right’ or ‘right-over-left’.

Cross References
Cover tests; Heterotropia; Hypertropia
Ice Pack Test
The ice pack test, or ice-on-eyes test, is performed by holding an ice cube, wrapped in a towel or a surgical glove, over the levator palpebrae superioris muscle of a ptotic eye for 2–10 min. Improvement of ptosis is said to be specific for myasthenia gravis, perhaps because cold improves transmission at the neuromuscular junction (myasthenic patients often improve in cold as opposed to hot weather). This phenomenon is generally not observed in other causes of ptosis, although it has been reported in Miller Fisher syndrome. A pooled analysis of several studies gave a test sensitivity of 89% and specificity of 100% with correspondingly high positive and negative likelihood ratios. The test is easy to perform and without side effects (cf. tensilon test). Whether the ice pack test is also applicable to myasthenic diplopia has yet to be determined: false positives have been documented.

References

Cross References
Diplopia; Fatigue; Ptosis

Ideational Apraxia
- see APRAXIA

Ideomotor Apraxia (IMA)
- see APRAXIA

Illusion
An illusion is a misinterpretation of a perception (cf. delusion, hallucination). Illusions occur in normal people when they are tired, inattentive, in conditions of poor illumination, or if there is sensory impairment. They also occur in disease states, such as delirium, and psychiatric disorders (affective disorders, schizophrenia). Examples of phenomena which may be labelled illusory include

- **Visual**: illusory visual spread, metamorphopsia, palinopsia, polyopia, teleopsia, Pufihrich phenomenon, visual alloaesthesia, visual perseveration;
- **Auditory**: palinacusis;
- **Vestibular**: vertigo.
Illusory Visual Spread
- see VISUAL PERSEVERATION

Imitation Behaviour
Imitation behaviour is the reproduction by the patient of gestures (echopraxia) and/or utterances (echolalia) made by the examiner in front of the patient; these ‘echophenomena’ are made by the patient without preliminary instructions to do so. They are consistent and have a compulsive quality to them, perhaps triggered by the equivocal nature of the situation. There may be accompanying primitive reflexes, particularly the grasp reflex, and sometimes utilization behaviour.

Imitation behaviour occurs with frontal lobe damage; originally mediobasal disease was thought the anatomical correlate, but more recent studies suggest upper medial and lateral frontal cortex. Certainly imitation behaviour never occurs with retrolateral cortical lesions.

A distinction has been drawn between ‘naïve’ imitation behaviour, which ceases after a direct instruction from the examiner not to imitate his/her gestures, which may be seen in some normal individuals; and ‘obstinate’ imitation behaviour which continues despite an instruction to stop; the latter is said to be exclusive to frontotemporal dementia.

References

Cross References
Echolalia; Echopraxia; Grasp reflex; Utilization behaviour

Imitation Synkinesis
- see MIRROR MOVEMENTS

Impersistence
Impersistence is an inability to sustain simple motor acts, such as conjugate gaze, eye closure, protrusion of the tongue, or keeping the mouth open. It is most commonly seen with lesions affecting the right hemisphere, especially central and frontal mesial regions, and may occur in association with left hemiplegia, neglect, anosognosia, hemianopia, and sensory loss. These patients may also manifest perseveration, echolalia, and echopraxia.
Impersistence is most often observed following vascular events but may also be seen in Alzheimer’s disease and frontal lobe dementias, and metabolic encephalopathies. Impersistence of tongue protrusion and handgrip may be seen in Huntington’s disease. Neuropsychologically, impersistence may be related to mechanisms of directed attention which are needed to sustain motor activity.

References

Cross References
Anosognosia, Echolalia; Echopraxia; Hemianopia; Milkmaid’s grip; Neglect; Perseveration; Trombone tongue

Inattention
- see NEGLECT

Incontinence
Urinary incontinence may result from neurological, as well as urological, disease. Neurological pathways subserving the appropriate control of micturition encompass the medial frontal lobes, a micturition centre in the dorsal tegmentum of the pons, spinal cord pathways, Onuf’s nucleus in the spinal cord segments S2–S4, the cauda equina, and the pudendal nerves. Thus, the anatomical differential diagnosis of neurological incontinence is broad. Moreover, incontinence may be due to inappropriate bladder emptying or a consequence of loss of awareness of bladder fullness with secondary overflow. Other features of the history and/or examination may give useful pointers as to localization. Incontinence of neurological origin is often accompanied by other neurological signs, especially if associated with spinal cord pathology (see Myelopathy). The pontine micturition centre lies close to the medial longitudinal fasciculus and local disease may cause an internuclear ophthalmoplegia. However, other signs may be absent in disease of the frontal lobe or cauda equina. Causes of urinary incontinence include

- Idiopathic generalized epilepsy with tonic–clonic seizures; however, the differential diagnosis of ‘loss of consciousness with incontinence’ also encompasses syncopal attacks with or without secondary anoxic convulsions, non-epileptic attacks, and hyperekplexia.
- Frontal lobe lesions: frontal lobe dementia; normal pressure hydrocephalus.
- Spinal cord pathways: urge incontinence of multiple sclerosis; loss of awareness of bladder fullness with retention of urine and overflow in tabes dorsalis.
- Sacral spinal cord injury; degeneration of the sacral anterior horn cells in Onuf’s nucleus (multiple system atrophy).
- Cauda equina syndrome; tethered cord syndrome (associated with spinal dysraphism).
- Pelvic floor injury.
Neurogenic incontinence may be associated with urgency, which results from associated abrupt increases in detrusor pressure (detrusor hyperreflexia); this may be helped by anticholinergic medication (e.g. oxybutynin, tolterodine). In addition there may be incomplete bladder emptying, which is usually asymptomatic, due to detrusor sphincter dyssynergy; for post-micturition residual volumes of greater than 100 ml (assessed by in–out catheterization or ultrasonography), this is best treated by clean intermittent self-catheterization.

References

Cross References
Cauda equina syndrome; Dementia; Frontal lobe syndromes; Hyperekplexia; Internuclear ophthalmoplegia; Myelopathy; Seizures; Urinary retention

Intention Myoclonus
- see MYOCLONUS

Intermanual Conflict
Intermanual conflict is a behaviour exhibited by an alien hand (le main étranger) in which it reaches across involuntarily to interfere with the voluntary activities of the contralateral (normal) hand. Diagonistic dyspraxia probably refers to the same phenomenon. The hand acts at cross purposes to the other following voluntary activity. A ‘compulsive grasping hand’ syndrome has been described which may be related to intermanual conflict, the difference being grasping of the contralateral hand in response to voluntary movement.

Intermanual conflict is more characteristic of the callosal, rather than the frontal, subtype of anterior or motor alien hand. It is most often seen in patients with corticobasal degeneration, but may also occur in association with callosal infarcts or tumours or following callosotomy.

Cross References
Alien hand, Alien limb; ‘Compulsive grasping hand’; Diagonistic dyspraxia

Intermetamorphosis
A form of delusional misidentification in which people known to the patient are believed to exchange identities with each other (cf. Fregoli syndrome, in which one person can assume different physical appearance).

Reference

Cross Reference
Delusion

Internal Ophthalmoplegia
- see OPHTHALMOPARESIS, OPHTHALMOPLEgia
Internuclear Ophthalmoplegia (INO)

Internuclear ophthalmoplegia (INO), or medial longitudinal fasciculus syndrome, consists of ipsilateral weakness of eye adduction with contralateral nystagmus of the adducting eye (ataxic or dissociated nystagmus), but with preserved convergence. This may be obvious with pursuit eye movements, but is better seen when testing reflexive saccades or optokinetic responses when the adducting eye is seen to ‘lag’ behind the adducting eye. INO may be asymptomatic or, rarely, may cause diplopia, oscillopsia, or a skew deviation. INO may be unilateral or bilateral. The eyes are generally aligned in primary gaze, but if there is associated exotropia this may be labelled wall-eyed monocular or bilateral internuclear ophthalmoplegia (WEMINO, WEBINO syndromes).

The pathoanatomical substrate of INO is pathology affecting the medial longitudinal fasciculus, a pathway linking the nuclei of cranial ocular motor nerves (III, IV, VI). The most common cause of INO by far is demyelination, particularly in young patients, but other causes include infarction (particularly older patients), Wernicke–Korsakoff syndrome, infection, trauma, tentorial herniation, haemorrhage, vasculitis, and paraneoplasia.

A similar clinical picture may be observed with pathology elsewhere, hence a ‘false-localizing’ sign and referred to as a pseudointernuclear ophthalmoplegia, especially in myasthenia gravis.

References

Cross References
Diplopia; ‘False-localizing signs’; One-and-a-half syndrome; Optokinetic nystagmus, Optokinetic response; Oscillopsia; Pseudointernuclear ophthalmoplegia; Saccades; Skew deviation

Intrusion

An intrusion is an inappropriate recurrence of a response (verbal, motor) to a preceding test or procedure after intervening stimuli. Schnider characterizes them as a form of confabulation. Intrusions are thought to reflect inattention and may be seen in dementing disorders or delirium. These phenomena overlap to some extent with the recurrent type of perseveration.

The term intrusion is also used to describe inappropriate saccadic eye movements which interfere with macular fixation during pursuit eye movements.

References

Cross References
Confabulation; Delirium; Dementia; Perseveration; Saccadic intrusion, Saccadic pursuit
Inverse Marcus Gunn Phenomenon
- see JAW WINKING; PTOSIS

Inverse Uhthoff Sign
- see UHTHOFF’S PHENOMENON

Inverted Reflexes
A phasic tendon stretch reflex is said to be inverted when the movement elicited is opposite to that normally seen, e.g. extension of the elbow rather than flexion when eliciting the supinator (brachioradialis) jerk; flexion of the forearm when tapping the triceps tendon (paradoxical triceps reflex); and flexion (hamstring contraction) rather than extension of the knee when tapping the patellar tendon.

The finding of inverted reflexes may reflect dual pathology, but more usually reflects a single lesion which simultaneously affects a root or roots, interrupting the local reflex arc, and the spinal cord, damaging corticospinal (pyramidal tract) pathways which supply segments below the reflex arc. Hence, an inverted supinator jerk is indicative of a lesion at C5/6, paradoxical triceps reflex occurs with C7 lesions; and an inverted knee jerk indicates interruption of the L2/3/4 reflex arcs, with concurrent damage to pathways descending to levels below these segments.

Reference

Cross Reference
Reflexes

Ipsipulsion
- see LATEROPULSION

Iridoplegia
Paralysis of the iris, due to loss of pupillary reflexes. This may be partial, as in Argyll Robertson pupil or Holmes–Adie pupil, or complete as in the internal ophthalmoplegia of an oculomotor (III) nerve palsy.

Cross References
Argyll Robertson pupil; Holmes–Adie pupil, Holmes–Adie syndrome; Oculomotor (III) nerve palsy; Ophthalmoparesis, Ophthalmoplegia; Pupillary reflexes
Jacksonian March
Jacksonian march is the sequential spread of a simple partial seizure to involve other body parts, for example, jerking may spread from one hand up the arm, to the ipsilateral side of the face. It may culminate in a secondary generalized seizure. The pathophysiological implication is of electrical disturbance spreading through the homunculus of the motor cortex. A sensory equivalent occurs but is rare.

References

Cross Reference
Seizures

Jactitation
Jactitation is literally ‘throwing about’, but may also imply restlessness. The term has been used in various ways: to refer to jerking or convulsion of epileptic origin; or jerking of choreic origin; or of myoclonic origin, such as ‘hypnagogic jactitation’ (physiological myoclonus associated with falling to sleep). It may also be used to refer to the restlessness seen in acute illness, high fever, and exhaustion, though differing from the restlessness implied by akathisia. Hence, it is essentially a non-specific term.

Cross References
Akathisia; Myoclonus; Seizures

Jamais Entendu
A sensation of unfamiliarity akin to jamais vu but referring to auditory experiences.

Jamais Vécu
- see JAMAIS VU

Jamais Vu
Jamais vu (literally ‘never seen’) and jamais vécu (‘never lived’) are complex auras of focal onset epilepsy in which there is a sensation of strangeness or unfamiliarity about visual stimuli that have in fact been previously experienced (cf. déjà vu). This is suggestive of seizure onset in the limbic system, but is not lateralizing (cf. déjà vu).

Cross References
Aura; Déjà vu
Jargon Aphasia

Jargon aphasia is a fluent aphasia characterized by a jumbled, unintelligible and meaningless (to the listener) output, with multiple paraphasias and neologisms, and sometimes echolalia (as in transcortical sensory aphasia). There may be a pressure of speech (logorrhoea). There is debate as to whether jargon aphasia is simply a primary Wernicke/posterior/sensory type of aphasia with failure to self-monitor speech output, or whether additional deficits (e.g. pure word deafness, intellectual impairment) are also required. Others suggest that jargon aphasia represents aphasia and anosognosia, leading to confabulation and reduplicative paramnesia.

References


Cross References

Anosognosia; Aphasia; Confabulation; Echolalia; Logorrhoea; Pure word deafness; Reduplicative paramnesia; Transcortical aphasias; Wernicke’s aphasia

Jaw Jerk

The jaw jerk, or masseter reflex, is contraction of the masseter and temporalis muscles in response to a tap on the jaw with the mouth held slightly open. Both the afferent and efferent limbs of the arc run in the mandibular division of the trigeminal (V) nerve, connecting centrally with the mesencephalic (motor) nucleus of the trigeminal nerve. The reflex is highly reproducible; there is a linear correlation between age and reflex latency and a negative correlation between age and reflex amplitude.

Interruption of the reflex arc leads to a diminished or absent jaw jerk as in bulbar palsy (although an absent jaw jerk may be a normal finding, particularly in the elderly). Bilateral supranuclear lesions cause a brisk jaw jerk, as in pseudobulbar palsy (e.g. in motor neurone disease).

Reference


Cross References

Age-related signs; Bulbar palsy; Pseudobulbar palsy; Reflexes

Jaw Winking

Jaw winking, also known as the Marcus Gunn phenomenon, is widening of a congenital ptosis when a patient is chewing, swallowing, or opening the jaw (i.e. a trigemino-oculomotor synkinesis). It is believed to result from aberrant innervation of the pterygoid muscles and levator palpebrae superioris.

Eyelid closure on jaw movement or opening of the mouth, the inverse Marcus Gunn phenomenon, is also described, as the Marin-Amat syndrome, thought to be due to aberrant facial (VII) nerve regeneration.
Jugular Foramen Syndrome

Reference

Cross References
Ptosis; Synkinesis, Synkinesia

**Jendrassik’s Manoeuvre**
Jendrassik’s manoeuvre is used to enhance or bring out absent or depressed tendon (phasic stretch) reflexes by isometric contraction of distant muscle groups, e.g. clenching teeth, or making a fist, interlocking fingers, and pulling the hands against one another. If previously absent reflexes are then elicited, this may be denoted +/−. Cocontraction increases the gain in the monosynaptic reflex arc, as distinct from facilitation or posttetanic potentiation which is seen in Lambert–Eaton myasthenic syndrome following tetanic contraction of muscles involved in the reflex.

References
Jendrassik E. Ueber allgemeine Localisation der Reflexe. *Deutsche Archiv fur Klinische Medicin* 1894; **52**: 569–600.

Cross References
Facilitation; Reflexes

**Jitteriness**
Jitteriness implies an exaggerated startle response, reflecting CNS overactivity. This may be confused in neonates with clonic seizures, but in the former there is stimulus sensitivity and an absence of associated ocular movements. However, both may occur in hypoxic–ischaemic or metabolic encephalopathies or with drug withdrawal.

Cross Reference
Seizures

**Joint Position Sense**
- see PROPRIOCEPTION

**Jugular Foramen Syndrome**
The glossopharyngeal (IX), vagus (X), and accessory (XI) cranial nerves may be damaged by lesions at or around the jugular foramen, producing a jugular foramen (or Vernet’s) syndrome. This produces

- Dysphagia, dysphonia, palatal droop, impaired gag reflex; ipsilateral reduced taste sensation on the posterior one-third of the tongue, and anaesthesia of the posterior one-third of the tongue, soft palate, pharynx, larynx, and uvula, due to glossopharyngeal and vagus nerve involvement.
- Ipsilateral weakness and atrophy of sternocleidomastoid and trapezius due to accessory nerve involvement (atrophy may be the more evident, hence the importance of palpating the muscle bellies).
Recognized causes of the jugular foramen syndrome include

- Skull base trauma/fracture;
- Glomus jugulare tumour;
- Inflammatory/infective collection at the skull base;
- Ischaemia.

The differential diagnosis includes retropharyngeal or retroparotid space occupying lesions, which may in addition involve the hypoglossal nerve (XII; Collet–Sicard syndrome) and the sympathetic chain with or without the facial nerve (VII; Villaret’s syndrome).

Cross References
Dysphagia; Dysphonia; Gag reflex

**Junctional Scotoma, Junctional Scotoma of Traquair**

Despite the similarity of these terms, they are used to refer to different types of scotoma:

- **Junctional scotoma:**
  
  Unilateral central scotoma with contralateral superior temporal defect, seen with lesions at the anterior angle of the chiasm. Such lesions have been said to damage the ipsilateral optic nerve plus the crossing loop of fibres (Wilbrand’s knee) originating from the inferonasal portion of the contralateral eye, although it may be noted that some authors have questioned whether such a loop in fact exists.

- **Junctional scotoma of Traquair:**
  
  A monocular temporal scotoma, sometimes even hemianopia, seen with optic nerve involvement sufficiently close to the chiasm to involve only ipsilateral crossing nasal axons, which subserve the temporal visual field, but sparing nasal axons crossing from the contralateral eye.

Reference

Cross References
Scotoma; Visual field defects
Kayser–Fleischer Rings

Kayser–Fleischer rings are deposits of copper, seen as a brownish discoloration, in Descemet’s membrane. Although often visible to the naked eye (difficult in people with a brown iris), they are best seen with slit-lamp examination. Since they are a highly reliable sign of intracerebral copper deposition in Wilson’s disease (hepatolenticular degeneration), any patient suspected of this diagnosis (i.e. with parkinsonism or dystonia presenting before age 50 years) should have a slit-lamp examination (as well as blood copper and caeruloplasmin, and urinary copper, measurements). Very occasionally cases of neurological Wilson’s disease without Kayser–Fleischer rings have been reported.

Reference

Cross References
Dystonia; Parkinsonism

Kernig’s Sign

Kernig’s sign is pain in the lower back (and also sometimes the neck) and resistance to movement with passive extension of the knee on the flexed thigh in a recumbent patient. It is indicative of meningeal mechanosensitivity due to inflammation, either infective (meningitis) or chemical (subarachnoid haemorrhage), in which case it may coexist with nuchal rigidity and Brudzinski’s (neck) sign. If unilateral it may indicate irritation of the lumbosacral nerve roots from a ruptured intervertebral disc (in which case Lasègue’s sign may also be present).

Reference

Cross References
Brudzinski’s (neck) sign; Lasègue’s sign; Nuchal rigidity

Kernohan’s Notch Syndrome

Raised intracranial pressure as a result of an expanding supratentorial lesion (e.g. tumour, subdural haematoma) may cause herniation of brain tissue through the tentorium into the subtentorial space, putting pressure on the midbrain. If the midbrain is shifted against the contralateral margin (free edge) of the tentorium, the cerebral peduncle on that side may be compressed, resulting in a hemiparesis which is ipsilateral to the supratentorial lesion (and hence may be considered ‘false-localizing’). There may also be an oculomotor nerve palsy ipsilateral to the lesion, which may be partial (unilateral pupil dilatation).
Kinesis Paradoxa

Kinesis paradoxica is the brief but remarkably rapid and effective movement sometimes observed in patients with Parkinson’s disease or postencephalitic parkinsonism, despite the poverty and slowness of spontaneous movement (akinesia, hypokinesia; bradykinesia) seen in these conditions. It often occurs in response to alarm, excitement, or emotion (e.g. in response to a genuinely funny joke).

Cross References
Akinesia; Bradykinesia; Hypokinesia; Parkinsonism

Klazomania

Klazomania was the term applied to the motor and vocal tics seen as a sequel to encephalitis lethargica (von Economo’s disease), along with parkinsonism and oculogyric crises. This observation helped to promote the idea that tics were due to neurological disease rather than being psychogenic, for example, in Tourette syndrome.

Reference

Cross References
Coprolalia; Echolalia; Parakinesia, Parakinesis; Tic

Kleptomania

Kleptomania, a morbid impulse to steal, has been related to the obsessive–compulsive spectrum of behaviours in patients with frontal lobe dysfunction.

Reference

Cross Reference
Frontal lobe syndromes

Klüver–Bucy Syndrome

The Klüver–Bucy syndrome consists of a variety of neurobehavioural changes, originally observed following bilateral temporal lobectomy (especially anterior tip) in monkeys, but subsequently described in man. The characteristic features, some or all of which may be present, are as follows:
Knee Tremor

- Visual agnosia (e.g. misrecognition of others)
- Hyperorality
- Hyperphagia, binge eating
- Hypermetamorphosis
- Hypersexuality
- Emotional changes: apathy; loss of fear, rage reactions

Clinical causes of the Klüver–Bucy syndrome include

- Sequel of bilateral temporal lobectomy
- Postictal phenomenon in a patient with a previous unilateral temporal lobectomy
- Sequel to minor head trauma; subdural haematoma
- Tumour
- Meningoencephalitis
- Pick’s disease
- Alzheimer’s disease: especially hyperorality and hyperphagia, but it is rare to have all features

References

Cross References
Apathy; Hypermetamorphosis; Hyperorality; Hyperphagia; Hypersexuality; Visual agnosia

Knee Tremor

A characteristic tremor of the patellae, sometimes known as knee bobbing, judder ing, or quivering, may be seen in primary orthostatic tremor (POT; ‘shaky legs syndrome’, ‘White rabbit syndrome’). It is due to rapid rhythmic contractions of the leg muscles on standing, which dampen or subside on walking, leaning against a wall, or being lifted off the ground, with disappearance of the knee tremor; hence this is a task-specific tremor. Auscultation with the diaphragm of a stethoscope over the lower limb muscles reveals a regular thumping sound, likened to the sound of a distant helicopter. EMG studies show pathognomonic synchronous activity in the leg muscles with a frequency of 14–18 Hz, thought to be generated by a central oscillator (peripheral loading does not alter tremor frequency).

A number of drugs have anecdotally been reported to be helpful in POT, including phenobarbitone, primidone, clonazepam, pramipexole, and levodopa, although the only blinded placebo-controlled study suggesting efficacy is with gabapentin. Unlike the situation in essential tremor, propranolol is not helpful.

References

Cross Reference
Tremor
Körber–Salus–Elschnig Syndrome
This describes convergence–retraction nystagmus, in which adducting saccades (medial rectus contraction) occur spontaneously or on attempted upgaze, often accompanied by retraction of the eyes into the orbits. This is associated with mesencephalic lesions of the pretectal region (e.g. pinealoma). The term may be used interchangeably with Parinaud’s syndrome or pretectal syndrome.

Cross References
Nystagmus; Parinaud’s syndrome

Kyphoscoliosis
Kyphoscoliosis is twisting of the spinal column in both the anteroposterior (kyphosis) and lateral (scoliosis) planes. Although such deformity is often primary or idiopathic, thus falling within the orthopaedic field of expertise, it may also be a consequence of neurological disease which causes weakness of paraspinal muscles. Recognized neurological associations of kyphoscoliosis and scoliosis include

- Chiari I malformation, syringomyelia
- Myelopathy (cause or effect? Skeletal disease such as achondroplasia is more likely to be associated with myelopathy than idiopathic scoliosis)
- Cerebral palsy
- Friedreich’s ataxia
- Neurofibromatosis
- Hereditary motor and sensory neuropathies
- Spinal muscular atrophies
- Myopathies, e.g. Duchenne muscular dystrophy

Stiff person syndrome may produce a characteristic hyperlordotic spine. Some degree of scoliosis occurs in virtually all patients who suffer from paralytic poliomyelitis before the pubertal growth spurt.

Cross References
Camptocormia; Stiffness
Lagophthalmos
Lagophthalmos is an inability to close the eyelid in a peripheral facial (VII) nerve palsy, with partial opening of the palpebral fissure. A similar phenomenon may be observed with aberrant regeneration of the oculomotor nerve, thought to be due to cocontraction of the levator palpebrae superioris and superior rectus muscles during Bell’s phenomenon.

Cross References
Bell’s palsy; Bell’s phenomenon; Facial paresis, Facial weakness

Lambert’s Sign
Lambert’s sign is a gradual increase in force over a few seconds when a patient with Lambert–Eaton myasthenic syndrome is asked to squeeze the examiner’s hand as hard as possible, reflecting increased power with sustained exercise. This may also be known as augmentation.

Cross Reference
Facilitation

Lasègue’s Sign
Lasègue’s sign is pain along the course of the sciatic nerve induced by stretching of the nerve, achieved by flexing the thigh at the hip while the leg is extended at the knee (‘straight leg raising’). This is similar to the manoeuvre used in Kernig’s sign (gradual extension of knee with thigh flexed at hip). Both indicate irritation of the lower lumbosacral nerve roots and/or meninges. The test may be positive with disc protrusion, intraspinal tumour, or inflammatory radiculopathy. Pain may be aggravated or elicited sooner using Bragard’s test, dorsiflexing the foot while raising the leg thus increasing sciatic nerve stretch, or Neri’s test, flexing the neck to bring the head on to the chest, indicating dural irritation.

A positive straight leg raising test is reported to be a sensitive indicator of nerve root irritation, proving positive in 95% of those with surgically proven disc herniation. The specificity may be somewhat lower.

Various modifications of Lasègue’s sign have been described. Crossed straight leg raising, when the complaint of pain on the affected side occurs with raising of the contralateral leg, is said to be less sensitive but highly specific. Femoral stretch test or ‘reverse straight leg raising’ may detect L3 root or femoral nerve irritation.

Reference

Cross References
Femoral stretch test; Kernig’s sign
Lateral Medullary Syndrome

The lateral medullary syndrome (or Wallenberg’s syndrome, after the neurologist who described it in 1895) results from damage (usually infarction) of the posterolateral medulla with or without involvement of the inferior cerebellum, producing the following clinical features:

- Nausea, vomiting, vertigo, oscillopsia (involvement of vestibular nuclei);
- Contralateral hypoalgesia, thermoanaesthesia (spinothalamic tract);
- Ipsilateral facial hypoalgesia, thermoanaesthesia, + facial pain (trigeminal spinal nucleus and tract);
- Horner’s syndrome (descending sympathetic tract), +/- ipsilateral hypo-hidrosis of the body;
- Ipsilateral ataxia of limbs (olivocerebellar/spinocerebellar fibres, inferior cerebellum);
- Dysphagia, dysphonia, impaired gag reflex;
- +/- eye movement disorders, including nystagmus, abnormalities of ocular alignment (skew deviation, ocular tilt reaction, environmental tilt), smooth pursuit and gaze holding, and saccades (lateropulsion);
- +/- hiccups (singultus); loss of sneezing.

Infarction due to vertebral artery occlusion (occasionally posterior inferior cerebellar artery) or dissection is the most common cause of lateral medullary syndrome, although tumour, demyelination, and trauma are also recognized causes.

References

Cross References
Anaesthesia; Dysphagia; Dysphonia; Environmental tilt; Gag reflex; Hemiataxia; Hiccups; Horner’s syndrome; Hypoalgesia; Hypohidrosis; Medial medullary syndrome; Nystagmus; Ocular tilt reaction; Oscillopsia; Saccades; Skew deviation; Sneezing; Vertigo

Lateral Rectus Palsy
- see ABDUCENS (VI) NERVE PALSY

Laterocollis
Laterocollis is a lateral head tilt; this may be seen in 10–15% of patients with torticollis.

Cross Reference
Torticollis
**Lateropulsion**
Lateropulsion or ipsipulsion is literally pulling to one side. The term may be used to describe ipsilateral axial lateropulsion after cerebellar infarcts preventing patients from standing upright causing them to lean towards the opposite side. Lateral medullary syndrome may be associated with lateropulsion of the eye towards the involved medulla, and there may also be lateropulsion of saccadic eye movements.

**Laughter**
- see AUTOMATISM; PATHOLOGICAL CRYING, PATHOLOGICAL LAUGHTER

**Lazarus Sign**
Various spontaneous and reflex movements are described in brain death, the most dramatic of which has been labelled as the Lazarus sign, after Lazarus, raised from the dead by Christ (John 11:1–44). This spinal reflex manifests as flexion of the arms at the elbow, adduction of the shoulders, lifting of the arms, dystonic posturing of the hands, and crossing of the hands.

**References**

**Leadpipe Rigidity**
- see RIGIDITY

**Leucocoria**
Leucocoria is a white pupillary reflex, in contrast to the normal red reflex. Causes include retinoblastoma, retinal detachment, toxocara infection, congenital cataract, and benign retinal hypopigmentation.

**Levator Inhibition**
- see EYELID APRAXIA

**Levitation**
Spontaneous levitation may be displayed by an alien limb, more usually an arm than a leg, indicative of parietal lobe pathology. It is most often seen in corticobasal (ganglionic) degeneration, but a few cases with pathologically confirmed progressive supranuclear palsy have been reported.

**References**

**Cross Reference**
Alien hand, Alien limb
Lhermitte’s Sign

Lhermitte’s sign, or the ‘barber’s chair syndrome’, is a painless but unpleasant tingling or electric shock-like sensation in the back and spreading instantaneously down the arms and legs following neck flexion (active or passive). It is associated with pathology within the cervical spinal cord. Although most commonly encountered (and originally described) in multiple sclerosis, it is not pathognomonic of demyelination and has been described with other local pathologies such as:

- subacute combined degeneration of the cord (vitamin B₁₂ deficiency);
- nitrous oxide (N₂O) exposure;
- traumatic or compressive cervical myelopathy (e.g. cervical spondylotic myelopathy);
- epidural/subdural/intraparenchymal tumour;
- radiation myelitis;
- pyridoxine toxicity;
- inflammation, e.g. systemic lupus erythematosus, Behçet’s disease;
- cervical herpes zoster myelitis;
- cavernous angioma of the cervical cord.

Pathophysiologically, this movement-induced symptom may reflect the exquisite mechanosensitivity of axons which are demyelinated or damaged in some other way.

A ‘motor equivalent’ of Lhermitte’s sign, McArdle’s sign, has been described, as has ‘reverse Lhermitte’s sign’, a label applied either to the aforementioned symptoms occurring on neck extension, or in which neck flexion induces electrical shock-like sensation travelling from the feet upward.

References


Cross References

McArdle’s sign; Myelopathy

Lid Lag

Lid lag is present if a band of sclera is visible between the upper eyelid and the corneal limbus on attempted downgaze (cf. lid retraction), seen for example in thyroid eye disease (Von Graefe’s sign), progressive supranuclear palsy (Steele–Richardson–Olszewski syndrome), and Guillain–Barré syndrome.

Cross References

Lid retraction; Von Graefe’s sign
Light-Near (Pupillary) Dissociation (LND)

Lid Retraction

Lid retraction is present if a band of sclera is visible between the upper eyelid and the corneal limbus in the primary position (cf. lid lag). This should be distinguished from contralateral ptosis. Recognized causes of lid retraction include:

- **Overactivity of levator palpebrae superioris:**
  - Dorsal mesencephalic lesion (Collier’s sign)
  - Opposite to unilateral ptosis, e.g. in myasthenia gravis; retracted lid may fall when ptotic lid raised; frontalis overactivity usually evident
  - Paradoxical lid retraction with jaw movement (jaw winking, Marcus Gunn phenomenon)

- **Overactivity of Müller’s muscle:**
  - Irritative oculosympathetic lesions (Claude–Bernard syndrome)

- **Contracture of the levator muscle:**
  - Hyperthyroidism, Graves’ ophthalmopathy (Dalrymple’s sign): may be associated lid lag
  - Myotonic syndromes
  - Aberrant oculomotor (III) nerve regeneration (pseudo-Von Graefe’s sign)

- Cicatricial retraction of the lid, e.g. following trauma
- Hepatic disease (Summerskill’s sign)
- Guillain–Barré syndrome

Lower lid retraction may be congenital, or a sign of proptosis. Ectropion may also be seen with lower lid tumour or chalazion, trauma with scarring, and ageing.

Cross References

Collier’s sign; Contracture; Dalrymple’s sign; Jaw winking; Lid lag; Proptosis; Pseudo-Von Graefe’s sign; Ptosis; Stellwag’s sign; Sunset sign

Light-Near (Pupillary) Dissociation (LND)

Light-near pupillary dissociation refers to the loss of pupillary light reflexes, whilst the convergence–accommodation reaction is preserved (see Pupillary Reflexes). This dissociation may be seen in a variety of clinical circumstances:

- **Argyll Robertson pupil:** small irregular pupils with reduced reaction to light, typically seen in neurosyphilis; the absence of miosis and/or pupillary irregularity has been referred to as pseudo-Argyll Robertson pupil, which may occur with neurosarcoidosis, diabetes mellitus, and aberrant regeneration of the oculomotor (III) nerve.

- **Holmes–Adie pupil:** dilated pupil showing strong but slow reaction to accommodation but minimal reaction to light (tonic > phasic).

- **Parinaud’s syndrome (dorsal rostral midbrain syndrome):** due to a lesion at the level of the posterior commissure and characterized by vertical gaze palsy, lid retraction (Collier’s sign) or ptosis, and large regular pupils responding to accommodation but not light.
Reference

Cross References
Argyll Robertson pupil; Collier’s sign; Holmes–Adie pupil, Holmes–Adie syndrome; Lid retraction; Parinaud's syndrome; Pseudo-Argyll Robertson pupil; Pupillary reflexes

Light Reflex
- see PUPILLARY REFLEXES

Locked-In Syndrome
The locked-in syndrome results from deafferentation, such that a patient is awake, self-ventilating, and alert, but unable to speak or move; vertical eye movements and blinking are usually preserved, affording a channel for simple (yes/no) communication.

The most common cause of the locked-in syndrome is basilar artery thrombosis causing ventral pontine infarction (both pathological laughter and pathological crying have on occasion been reported to herald this event). Other pathologies include pontine haemorrhage and central pontine myelinolysis. Bilateral ventral midbrain and internal capsule infarcts can produce a similar picture. Generally this is irreversible, although recovery has on occasion been recorded.

The locked-in syndrome may be mistaken for abulia, akinetic mutism, coma, and catatonia.

References

Cross References
Abulia; Akinetic mutism; Blinking; Catatonia; Coma; Pathological crying, Pathological laughter

Lockjaw
- see TRISMUS

Logoclonia
Logoclonia is the tendency for a patient to repeat the final syllable of a word when speaking; hence it is one of the reiterative speech disorders (cf. echolalia, palilalia); Liepmann apparently coined this term in 1905 to indicate ‘continuous perseveration’. It may be described as the festinating repetition of individual phonemes.
Logoclonia is an indicator of bilateral brain injury, usually involving subcortical structures, and may be seen in the late stages of dementia of Alzheimer type (but not in semantic dementia).

**Reference**

**Cross References**
Echolalia; Festination, Festinant gait; Palilalia; Perseveration

**Logopenia**
Logopenia is a reduced rate of language production, due especially to word-finding pauses, but with relatively preserved phrase length and syntactically complete language, seen in aphasic syndromes, such as primary non-fluent aphasia.

**Cross Reference**
Aphasia

**Logorrhoea**
Logorrhoea is literally a flow of speech, or pressure of speech, denoting an excessive verbal output, an abnormal number of words produced during each utterance. Content is often irrelevant, disconnected, and difficult to interpret. The term may be used for the output in the Wernicke/posterior/sensory type of aphasia or for an output which superficially resembles Wernicke aphasia but in which syntax and morphology are intact, rhythm and articulation are usually normal, and paraphasias and neologisms are few. Moreover, comprehension is better than anticipated in the Wernicke type of aphasia. Patients may be unaware of their impaired output (anosognosia) due to a failure of self-monitoring.

Logorrhoea may be observed in subcortical (thalamic) aphasia, usually following recovery from lesions (usually haemorrhage) to the anterolateral nuclei. Similar speech output may be observed in psychiatric disorders such as mania and schizophrenia (schizophrenia).

**Reference**

**Cross References**
Aphasia; Delirium; Echolalia; Jargon aphasia; Schizophrenia; Wernicke’s aphasia

**Long Tract Signs**
- see UPPER MOTOR NEURONE (UMN) SYNDROME

**Looking Glass Syndrome**
- see MIRROR AGNOSIA

**Lower Motor Neurone (LMN) Syndrome**
A lower motor neurone (LMN) syndrome constitutes a constellation of motor signs resulting from damage to lower motor neurone pathways, i.e. from anterior horn cell distally, encompassing the motor roots, nerve plexuses, peripheral
nerves, and neuromuscular junction. Following the standard order of neurological examination of the motor system, the signs include

- **Appearance:**
  muscle wasting; fasciculations (or ‘fibrillations’) may be observed or induced, particularly if the pathology is at the level of the anterior horn cell.

- **Tone:**
  reduced tone (flaccidity, hypotonus), although this may simply reflect weakness.

- **Power:**
  weakness, often marked; depending on the precise pathological process, weakness often affects both flexor and extensor muscles equally (although this is not always the case).

- **Coordination:**
  depending on the degree of weakness, it may not be possible to comment on the integrity or otherwise of coordination in LMN syndromes; in a pure LMN syndrome coordination will be normal.

- **Reflexes:**
  depressed (hyporeflexia) or absent (areflexia); plantar responses are flexor.

It is often possible to draw a clinical distinction between motor symptoms resulting from lower or upper motor neurone pathology and hence to formulate a differential diagnosis and direct investigations accordingly. Sensory features may also be present in LMN syndromes if the pathology affects sensory as well as motor roots, or both motor and sensory fibres in peripheral nerves.

**Cross References**
Areflexia; Fasciculation; Fibrillation; Flaccidity; Hyporeflexia; Hypotonia, Hypotonus; Neuropathy; Reflexes; Upper motor neurone (UMN) syndrome; Weakness

**Luria Test**
- see FIST–EDGE–PALM TEST
Macrographia
Macrographia is abnormally large handwriting. It may be seen in cerebellar disease, possibly as a reflection of the kinetic tremor and/or the impaired checking response seen therein (cf. micrographia).

Reference

Cross References
Micrographia; Tremor

Macropsia
Macropsia, or ‘Brobdingnagian sight’, is an illusory phenomenon in which the size of a normally recognized object is overestimated.

Cross Reference
Metamorphopsia

MacroSomatognosia
- see ‘ALICE IN WONDERLAND’ SYNDROME

MacroSquare Wave Jerks
- see SQUARE WAVE JERKS

Macula Sparing, Macula Splitting
Macula sparing is a feature of a homonymous hemianopia in which central vision is intact, due to damage confined to the occipital cortex without involving the occipital pole. This may occur because anastomoses between the middle and posterior cerebral arteries maintain that part of area 17 necessary for central vision after occlusion of the posterior cerebral artery.

Cortical blindness due to bilateral (sequential or simultaneous) posterior cerebral artery occlusion may leave a small central field around the fixation point intact, also known as macula sparing.

Macula splitting, a homonymous hemianopia which cuts through the vertical meridian of the macula, occurs with lesions of the optic radiation.

Hence, macula sparing and macula splitting have localizing value when assessing homonymous hemianopia.

Cross References
Cortical blindness; Hemianopia; Visual field defects
Maculopathy

Maculopathy is any process affecting the macula, with changes observable on ophthalmoscopy. These processes may produce a central or ring scotoma and visual failure. Common causes include

- *Diabetes mellitus*: oedema and hard exudates at the macula are a common cause of visual impairment, especially in non-insulin-dependent diabetes mellitus.
- *Hypertension*: abnormal vascular permeability around the fovea may produce a macular star.
- *Drug-induced*: e.g. ‘bull’s-eye’ maculopathy of chloroquine.
- ‘Cherry red spot at the macula’: this appearance may occur in sialidosis (‘cherry red spot–myoclonus syndrome’) and gangliosidoses (e.g. Tay–Sachs disease).

**Cross References**
Cherry red spot at the macula; Retinopathy; Scotoma; Visual field defects

Magnetic Movements

Movements may be described as magnetic in varying contexts:

- the following or tracking movements of an alien hand in corticobasal degeneration, reaching out to touch or grasp the examiner's hand or clothing, as in forced groping; also known as compulsive tactile exploration;
- in a hesitant gait (ignition failure), with seeming inability to lift the feet (‘stuck to the floor’) in gait apraxia.

**Reference**

**Cross References**
Alien hand, Alien limb; Forced groping; Gait apraxia; Grasp reflex

Main d'accoucheur

*Main d'accoucheur*, or carpopedal spasm, is a posture of the hand with wrist flexion in which the muscles are rigid and painful. *Main d'accoucheur* is so called because of its resemblance to the posture of the hand adopted for the manual delivery of a baby (‘obstetrical hand’).

This tetanic posture may develop in acute hypocalcaemia (induced by hyperventilation, for instance) or hypomagnesaemia and reflects muscle hyperexcitability. Development of *main d'accoucheur* within 4 min of inflation of a sphygmomanometer cuff above arterial pressure (Trousseau’s sign) indicates latent tetany. Mechanosensitivity of nerves may also be present elsewhere (Chvostek’s sign).

**Cross References**
Chvostek’s sign; Trousseau’s sign

Main en Griffe

- see CLAW HAND
Main Étranger
- see ALIEN HAND, ALIEN LIMB

Main Succulente
Main succulente refers to a swollen hand with thickened subcutaneous tissues, hyperkeratosis and cyanosis, and trophic changes which may be observed in an analgesic hand, e.g. in syringomyelia.

Cross Reference
Charcot joint

Man-in-a-Barrel
'Man-in-a-barrel' is a clinical syndrome of brachial diplegia with preservation of brainstem function and of muscle strength in the legs.

This most usually occurs as a result of bilateral borderzone infarcts in the territories between the anterior and middle cerebral arteries ('watershed infarction'). This may be as a consequence of cerebral hypoperfusion (e.g. during cardiac arrest, cardiac surgery), in which case the prognosis is poor. The clinical picture has also been reported with cerebral metastases. Acute central cervical cord lesions may also produce a ‘man-in-a-barrel’ syndrome, for example, after severe hyperextension injury or after unilateral vertebral artery dissection causing anterior cervical spinal cord infarction. This may follow a transient quadriplegia, and considerable recovery is possible.

A neurogenic man-in-a-barrel syndrome has been reported (‘flail arm syndrome’), which is a variant of motor neurone disease. Likewise, bilateral neuralgic amyotrophy can produce an acute peripheral man-in-a-barrel phenotype.

References

Cross References
Flail arm; Quadriparesis, Quadriplegia

Marche à Petit Pas
Marche à petit pas is a disorder of gait characterized by impairments of balance, gait ignition, and locomotion. Particularly there is shortened stride (literally marche à petit pas) and a variably wide base. This gait disorder is often associated with dementia, frontal release signs, and urinary incontinence, and sometimes with apraxia, parkinsonism, and pyramidal signs. This constellation of clinical signs reflects underlying pathology in the frontal lobe and subjacent white matter, most usually of vascular origin, and is often associated with a subcortical vascular dementia. Modern clinical classifications of gait disorders have subsumed marche à petit pas into the category of frontal gait disorder.
Reference

Cross References
Apraxia; Dementia; Frontal release signs; Parkinsonism

Marcus Gunn Phenomenon
- see JAW WINKING

Marcus Gunn Pupil, Marcus Gunn Sign
The Marcus Gunn pupil or sign, first described in 1902 by the ophthalmologist Robert Marcus Gunn, is the adaptation of the pupillary light reflex to persistent light stimulation, that is, a dilatation of the pupil is observed with continuing stimulation with incident light (‘dynamic anisocoria’). This is indicative of an afferent pathway defect, such as retrobulbar neuritis. The swinging flashlight sign or test may be used to demonstrate this by comparing direct and consensual pupillary light reflexes in one eye. Normally the responses are equal but in the presence of an afferent conduction defect an inequality is manifest as pupillary dilatation.

References

Cross References
Pupillary reflexes; Relative afferent pupillary defect (RAPD); Swinging flashlight sign

Mask-Like Facies
The poverty of spontaneous facial expression, hypomimia, seen in extrapyramidal disorders such as idiopathic Parkinson’s disease is sometimes described as mask-like.

Cross References
Hypomimia; Parkinsonism

Masseter Hypertrophy
Masseter hypertrophy, either unilateral or bilateral, may occur in individuals prone to bruxism. A familial syndrome of hypertrophy of the masseter muscles has been described.

Reference

Cross Reference
Bruxism

Masseter Reflex
- see JAW JERK
Masticatory Claudication
Pain in the muscles of mastication with chewing may be a sign, along with headache, of giant cell (temporal) arteritis.

McArdle’s Sign
McArdle’s sign is the combination of reduced lower limb strength, increased lower limb stiffness, and impaired mobility following neck flexion. The difference may best be appreciated by comparing leg strength (e.g. hip flexion) with the neck fully extended and fully flexed.

The sign was initially described in multiple sclerosis but may occur in other myelopathies affecting the cord at any point between the foramen magnum and the lower thoracic region. The mechanism is presumed to be stretch-induced conduction block, due to demyelinated plaques or other pathologies, in the corticospinal tracts. McArdle’s sign may be envisaged as the motor equivalent of Lhermitte’s sign.

References

Cross References
Lhermitte’s sign; Myelopathy

Medial Medullary Syndrome
The medial medullary syndrome, or Dejerine’s anterior bulbar syndrome, results from damage to the medial medulla, most usually infarction as a consequence of anterior spinal artery or vertebral artery occlusion. The clinical picture is of:

- Ipsilateral tongue paresis and atrophy, fasciculations (hypoglossal nerve involvement);
- Contralateral hemiplegia with sparing of the face (pyramid);
- Contralateral loss of position and vibration sense (medial lemniscus) with pain and temperature sensation spared;
- +/- upbeat nystagmus (?nucleus intercalatus of Staderini).

References

Cross References
Fasciculation; Hemiplegia; Lateral medullary syndrome; Nystagmus

Menace Reflex
- see BLINK REFLEX
Meningism
Meningism (meningismus, nuchal rigidity) is a stiffness or discomfort on passive movement (especially flexion) of the neck in the presence of meningeal irritation (e.g. infective meningitis, subarachnoid haemorrhage). A number of other, eponymous, signs of meningeal irritation have been described, of which the best known are those of Kernig and Brudzinski.

Meningism is not synonymous with meningitis, since it may occur in acute systemic pyrexial illnesses (pneumonia, bronchitis), especially in children. Moreover, meningism may be absent despite the presence of meningitis in the elderly and those receiving immunosuppression.

Cross References
Brudzinski’s (neck) sign; Kernig’s sign; Nuchal rigidity

Metamorphopsia
Metamorphopsia is an illusory visual phenomenon characterized by objects appearing distorted or misshapen in form. As with neglect, these phenomena may be classified as object- or person-centred:

- **Object-centred:** affecting size and spatial relationships.
  - *Macropsia:* objects appear larger than normal.
  - *Micropsia:* objects appear smaller than normal.
  - *Pelopsia:* objects appear closer to the observer than actual.
  - *Porropsia:* objects appear farther away from the observer than actual.

- **Person-centred:**
  - *Microsomatognosia and macrosomatognosia:* body image appears smaller or larger than normal (‘Alice in Wonderland’ syndrome).

Metamorphopsias are often transient and episodic, occurring, for example, during migraine attacks, epileptic seizures, with psychotropic drug abuse, and following petechial intraparenchymal haemorrhages. Rarely, they are long-lasting or permanent, for example, following brain infarction (most commonly involving the occipito-parietal or temporoparietal cortex: lesions on the right are more likely than those on the left to give metamorphopsia) or tumours. Retinal disease causing displacement of photoreceptors may produce metamorphopsia: micropsia due to receptor separation in retinal oedema, macropsia due to receptor approximation in retinal scarring. Occasional cases of metamorphopsia have been reported with lesions of the optic chiasm, optic radiation, and retrosplenial region. Indeed, it seems that metamorphopsia may occur with pathology at any point along the visual pathway from retina to cortex. Differing patterns of metamorphopsia may assist with clinico-anatomical correlation:

- retinal lesions: ipsilateral monocular;
- chiasmal lesions: bitemporal;
- occipitoparietal lesions: contralateral homonymous.

Metamorphopsia may be associated with visual hallucinations.

The Amsler Chart Manual (test charts to determine the quality of central vision, by Prof. Dr. Marc Amsler of Zurich) includes charts to demonstrate metamorphopsia (numbers 5 and 6).
Micropsia

Reference

Cross References
‘Alice in Wonderland’ syndrome; Hallucination; Illusion; Macropsia; Micropsia; Pelopsia; Porropsia; Telopsia

Micrographia

Micrographia is small handwriting. It is most often recognized in association with the extrapyramidal features of idiopathic Parkinson’s disease (indeed it may be the presenting sign), but may occasionally occur with other parkinsonian syndromes (e.g. progressive supranuclear palsy [PSP]) or in isolation with focal lesions of the midbrain or basal ganglia.

In Parkinson’s disease, handwriting may initially be of normal size but then becomes progressively smaller, slower, and more illegible as writing proceeds, an example of parkinsonian fatigue, a gradual decline in the amplitude and speed of initiation of voluntary movements. Such ‘slow’ micrographia may be distinguished from ‘fast’ micrographia in which letters are small throughout although written at normal speed without fatigue, which may be seen in PSP or other pallidal pathologies.

There is a poor correlation between micrographia and the side, severity, or duration of classical parkinsonian features, and its response to levodopa preparations is very variable. These observations, along with reports of isolated micrographia with cortical lesions demonstrated by neuroimaging, suggest that the anatomical basis of micrographia may be at the level of the cortex (dominant parietal lobe) rather than the basal ganglia.

Micrographia has also been described following large right anterior cerebral artery infarcts and lacunar infarcts involving the putamen and genu of the internal capsule. Isolated micrographia has been reported with lenticular haematoma.

References


Cross References
Fast micrographia; Fatigue; Parkinsonism

Micropsia

Micropsia, or ‘Lilliput sight’, is an illusory phenomenon in which the size of a normally recognized object is underestimated. It is the most common form of metamorphopsia and is most often associated with lesions of the right temporoparietal cortex, although macular oedema and optic chiasm lesions may also cause micropsia. In migraine transient micropsia may occur. Hemimicropsia,
micropsia confined to one visual hemifield, has been recorded. The entirely subjective nature of the disorder may account for the relative rarity of reports.

References

Cross-Reference
Metamorphopsia

**Microsomatognosia**
- see ‘ALICE IN WONDERLAND’ SYNDROME

**Milkmaid’s Grip**
Milkmaid’s grip is the descriptive term applied to the inability to maintain a firm grip (e.g. of the examiner’s fingers), detected as an alternating squeezing and releasing (as required for successful milking by hand). Seen in Huntington’s disease, this may reflect a combination of chorea and motor impersistence.

Cross References
Chorea, Choreoathetosis; Impersistence; Trombone tongue

**Miosis**
Miosis is abnormal reduction in pupillary size, which may be unilateral or bilateral. Causes include

- Oculosympathetic paresis of whatever cause, e.g. Horner’s syndrome (unilateral), pontine haemorrhage (bilateral), early stages of central cephalic herniation (bilateral);
- Drug-induced: e.g. opiates (bilateral);
- Pupils tend to be small and reactive in metabolic-toxic encephalopathies (bilateral);
- ‘Senile miosis’ (bilateral): age-related.

If only one pupil appears small (anisocoria), it is important to distinguish miosis from contralateral mydriasis, when a different differential will apply.

Reference

Cross References
Age-related signs; Anisocoria; Argyll Robertson pupil; Horner’s syndrome; Mydriasis

**Mirror Agnosia**
Mirror agnosia, or the ‘looking glass syndrome’, is a phenomenon observed in patients with left hemispatial neglect as a result of right parietal lobe lesions. There is inability to point to objects seen in a mirror, with repeated reaching ‘into’ the mirror even when the actual location of the target is shown. In a milder
form, known as ‘mirror ataxia’, patients reach in the direction of the object but with increased errors of reach and grasp, suggesting that visual information is not adequately transformed into a body-centred frame of reference.

**References**


**Cross References**

Agnosia; Neglect

**Mirror Apraxia**

Patients with mirror apraxia presented with an object that can be seen only in a mirror, when asked to reach for the real object will reach for the virtual object in the mirror. Mirror apraxia results from right or left posterior parietal lobe lesions.

**Reference**


**Mirror Ataxia**

- see MIRROR AGNOSIA

**Mirror Dystonia**

- see WRITER’S CRAMP

**Mirror Hallucination**

- see AUTOSCOPY

**Mirror Movements**

Mirror movements are involuntary movements of one side of the body that accompany and ‘mirror’ (reflect) intentional movements on the opposite side of the body (also known as imitation synkinesis). They are usually symmetrical and most often seen when using distal muscles of the upper limb. Mirror movements are frequently present in young children but prevalence decreases with age. Persistence of mirror movements into adult life (‘congenital mirror movements’) is pathological, as is acquisition in adult life. These movements are uncommon after acquired brain lesions with no relationship to specific anatomical areas.

Congenital mirror movements are associated with skeletal developmental abnormalities, especially of the atlanto-occipital region, such as Klippel–Feil syndrome. They are also seen in 85% of patients with X-linked Kallmann syndrome (hypogonadotrophic hypogonadism and anosmia).

Acquired mirror movements have been described following thalamic lesions, and in association with spastic paraparesis, extrapyramidal disorders
(Parkinson’s disease, multiple system atrophy), Friedreich’s ataxia, phenylketonuria, and affecting hemiparetic limbs following stroke in young children.

There is some neurophysiological evidence from patients with X-linked Kallmann syndrome for the existence of an ipsilateral corticospinal pathway, consistent with other evidence that the congenital condition is primarily a disorder of axonal guidance during development. Concurrent activity within ipsilateral and contralateral corticospinal pathways may explain mirroring of movements. Alternatively, a failure of transcallosal inhibition, acquired at the time of myelination of these pathways, may contribute to the genesis of mirror movements. Loss of joint position sense following thalamic lesions may be of relevance. A deficit of sustained attention has also been postulated as the cause of mirror movements.

References

Cross References
Anosmia; Attention; Mirror writing; Proprioception; Synkinesia, Synkinesis

**Mirror Sign**

The term ‘mirror sign’ has been applied to the phenomenon of misrecognition of self as another when seen in a mirror. It may be classified with the delusional misidentification syndromes. This may occur in Alzheimer’s disease and frontotemporal dementia and is associated with impaired cognition, confabulation, and prefrontal dysfunction. It may lead to a patient complaint of an intruder or a stranger living in the house (‘phantom boarder’ syndrome). Failure to recognize oneself in a mirror may also be a dissociative symptom, a symptom of depersonalization.

Some authors believe ‘the phenomenon of the mirror’ to be an extreme example of prosopagnosia, but other studies have not found an association.

Reference

Cross References
Confabulation; Depersonalization; Misidentification syndromes; ‘Picture sign’; Prosopagnosia

**Mirror Writing**

As the name implies, mirror writing is a mirror image of normal writing, hence running from right to left, with characters back to front. This may occur spontaneously, apparently more often in left-handers, or in right-handers attempting to write with the left hand following left-sided brain injury (e.g. stroke).
Leonardo da Vinci (1452–1519) is the most celebrated mirror writer: it is possible his left-handedness, and hence mirror writing, followed an injury to his right hand. The author Lewis Carroll occasionally wrote mirror letters but these differ from his normal script, unlike the situation with Leonardo whose two scripts are faithful mirror images. Carroll’s letters may thus reflect not an inherent capacity but a contrivance, designed to amuse children who corresponded with him. The device was also used by the author Arthur Ransome in his 1939 novel Secret Water. Jane Austen wrote one letter (1817) to a young niece in which script runs from right to left but with word order reversed within words (i.e. not mirror writing).

Various neural mechanisms are proposed to explain mirror writing, including bilateral cerebral representation of language, motor programmes, or visual memory traces or engrams. The mechanisms may differ between a true mirror writer like Leonardo and someone performing the task for amusement like Carroll.

Double mirror writing (écriture en double miroir) is inverted top to bottom (i.e. script goes up the page, upside down) in addition to being mirror reversed.

The ability to read mirror reversed text as quickly as normally oriented text has been reported in some autistic individuals.

References

Misidentification Syndromes
These are defined as delusional conditions in which patients incorrectly identify and reduplicate people, places, objects, or events. Examples include

- Capgras syndrome; may be related to reduplicative paramnesia
- Fregoli syndrome
- Intermetamorphosis
- Phantom boarder sign
- Mirror sign

References

Cross References
Delusion; Intermetamorphosis; ‘Mirror sign’; Reduplicative paramnesia

Misoplegia
Misoplegia is a disorder of body schema in which there is active hatred of a paralyzed limb, with or without personification of the limb, and attempts to injure the paralyzed limb. It occurs with right parietal region injury (hence left-sided limbs most often involved) and may occur in conjunction with anosognosia, left hemispatial neglect, and (so called) constructional apraxia.

References

Cross References
Anosognosia; Apraxia; Hemiparesis; Hemiplegia; Neglect

Mitbewegungen
- see SYNKINESIA, SYNKINESIS

Mitgehen
An abnormality of induced movement, in which limb movement occurs in response to application of the slightest pressure despite the patient having been told to resist (German: to go too); a manifestation of negativism.

Cross Reference
Negativism

Mitmachen
A motor disorder in which the patient acquiesces to every passive movement of the body made by the examiner, but as soon as the examiner releases the body part, the patient returns it to the resting position.

Monoballismus
Monoballismus is ballism affecting a single limb.

Cross References
Ballism, Ballismus; Hemiballismus
Monochromatopsia
The author has seen a patient with a diagnosis of frontotemporal dementia who persistently and consistently complained that everything he saw was red, even though he was aware that they were not red, for example, his wife’s grey hair. His speech was fluent without paraphasia although impoverished in content, with recurrent themes repeated almost verbatim. He had mild oro-facial dyspraxia. There was no alexia. Confronted with objects of different colours, he was unable to point to them by colour since all appeared red to him. The features seem to be distinct from erythropsia (persistent) or phantom chromatopsia (normal visual acuity). The author proposes that this phenomenon might be termed ‘monochromatopsia’.

Cross References
Erythropsia; Phantom chromatopsia

Monomelia
- see MONOPARESIS, MONOPLEGIA

Mononeuritis Multiplex, Mononeuropathy Multiplex
- see NEUROPATHY

Mononeuropathy
- see NEUROPATHY

Monoparesis, Monoplegia
Monoparesis is weakness, monoplegia complete weakness (‘paralysis’), of a single limb. Monoparesis of the arm or leg of upper motor neurone type is usually cortical in origin, although may unusually arise from a cord lesion (leg more frequently than arm). Hoover’s sign and Babinski’s trunk–thigh test may be helpful in deciding whether monoparetic/monoplegic leg weakness is of non-organic origin, and the ‘arm drop’ or ‘face–hand test’ in arm weakness.

Peripheral disorders can sometimes present exclusively with single limb weakness, such as monomelic motor neurone disease (Hirayama disease), multifocal motor neuropathy with conduction block, and Guillain–Barré syndrome.

Cross References
‘Arm drop’; Babinski’s trunk–thigh test; Hemiparesis; Hoover’s sign

Monophasia
- see RECURRENT UTTERANCES

Monotonia
Monotonia is a restricted range of speech inflection, occurring with hypophonia as part of the hypokinetic dysarthria observed in parkinsonism.

Cross References
Dysarthria; Hypophonia; Parkinsonism

Moria
Moria is literally folly (as in Desiderius Erasmus’ Moriae Encomium of 1509, literally ‘praise of folly’). In clinical usage, the meaning overlaps not only with
that of emotional lability but has also been used in the context of pathological laughter.

**Cross References**

Emotionalism, Emotional lability; Pathological crying, Pathological laughter; Witzelsucht

**Motor Neglect**

Motor neglect is failure to move the contralesional limbs in the neglect syndrome, a more severe impairment than directional hypokinesia.

**Cross References**

Directional hypokinesia; Eastchester clapping sign; Neglect

**Moving Ear**

A focal dyskinesia characterized by ear movement has been described.

**Reference**


**Cross Reference**

Dyskinesia

**Multiplication of Images**

- see POLYOPIA

**Muscle Hypertrophy**

Muscle hypertrophy is muscle enlargement due to an increase in the size of its myofibrils. Muscle hypertrophy may be generalized or focal and occurs in response to repetitive voluntary contraction (physiological) or repetitive abnormal electrical activity (pathological, e.g. myotonia in Thomsen's disease; primary orthostatic tremor). Muscle enlargement may also result from replacement of myofibrils by other tissues such as fat or amyloid, a situation better described as pseudohypertrophy.

**Cross References**

Calf hypertrophy; Masseter hypertrophy; Myotonia

**Mutism**

Mutism is absence of speech output. This may be psychogenic, as in schizophrenia or affective disorders, with or without catatonia; or a consequence of neurological disease, for example:

- Akinetic mutism;
- Dementia syndromes, especially frontal lobe dementia, late stages of primary non-fluent aphasia;
- Encephalopathy (toxic/drug-induced/metabolic);
- Damage to Broca's area, supplementary motor area; severe pseudobulbar palsy, bilateral thalamic damage;
- Cerebellar mutism: rare, following midline cerebellar surgery in children. Thought to be due to dentatothalamocortical tract damage, bilateral oedema in cerebellar peduncles (rather than surgical trauma or infarction);
- Bilateral vocal cord paralysis (although this may be better termed aphonia);
- Autism.
In neurological disorders there may be difficulty initiating movements, completing motor sequences, or inhibition of appropriate responses.

**References**

**Cross References**
Aphasia; Aphonia

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**Myasthenic Snarl**
Patients with weakness of facial musculature as a consequence of myasthenia gravis may have a ‘transverse smile’, with lack of elevation of the corners of the mouth, or appear to snarl when asked to smile or laugh. This may give the impression that they seem peculiarly unamused by an examiner’s attempted wit-ticisms. These phenomena may be seen with other causes of facial weakness, such as facioscapulohumeral (FSH) dystrophy.

**Mydriasis**
Mydriasis is an abnormal dilatation of the pupil, either unilateral or bilateral. Causes include

- Oculoparasympathetic paresis, from lesions at the Edinger–Westphal nucleus or anywhere along the course of the oculomotor (III) nerve (usually unilateral);
- Tonic enlargement of the pupil (Holmes–Adie pupil, usually unilateral);
- Sympathomimetic drugs, e.g. adrenaline (usually bilateral);
- Later stages of central cephalic herniation;
- A syndrome of benign episodic unilateral mydriasis has been described, sometimes related to migraine.

If only one pupil appears large (anisocoria), it is important to distinguish mydriasis from contralateral miosis, when a different differential will apply (e.g. Horner’s syndrome).

**References**

**Cross References**
Anisocoria; Holmes–Adie pupil, Holmes–Adie syndrome; Horner’s syndrome; Hutchinson’s pupil; Miosis; Oculomotor (III) nerve palsy
Myelopathy

A myelopathy is a disorder of the spinal cord. Such disorders may be further characterized according to whether the responsible lesion lies within or outside the spinal cord: intrinsic or intramedullary lesions are always intradural; extrinsic or extramedullary lesions may be intradural or extradural. It may be possible to differentiate intramedullary from extramedullary lesions on clinical grounds, although this distinction is never absolute because of clinical overlap.

Clinical features of extrinsic/extramedullary myelopathy:

- **Motor**: sequential spastic paraparesis below the level of the lesion; upper motor neurone (UMN) signs occur early; lower motor neurone (LMN) signs are unusual and have a segmental (radicular) distribution if present.

- **Sensory**: symptoms of pain may be radicular (e.g. secondary to a neurofibroma) or vertebral (e.g. secondary to neoplastic or inflammatory processes); sensory signs are not usually marked until the later stages, and all modalities are often involved. A Brown–Séquard syndrome may be more common in extrinsic than intrinsic myelopathies.

- **Sphincters**: may have bladder urgency, impotence.

Pathologies commonly causing extrinsic myelopathy include:

- prolapsed disc, osteophyte bar;
- tumour (primary, secondary);
- arteriovenous malformation/haematoma;
- abscess.

Clinical features of intrinsic/intramedullary myelopathy:

- **Motor**: LMN signs may be prominent and diffuse; UMN signs tend to occur late (spastic paraparesis below level of lesion). A combination of UMN and LMN signs is much more likely to reflect intrinsic than extrinsic pathology.

- **Sensory**: symptoms of central (funicular) pain may occur; dissociated sensory loss (spinothalamic > dorsal column involvement, or vice versa), suspended sensory loss, and sacral sparing are characteristic of intramedullary lesions; a Brown–Séquard syndrome may occur. Vibratory sensibility is more often affected than proprioception.

- **Sphincters**: bladder involvement common, often early and slow to recover.

These features are dependent on the extent to which the cord is involved: some pathologies have a predilection for posterior columns, central cord, etc. Pathologies commonly causing intrinsic myelopathy include:

- multiple sclerosis or other inflammatory process causing transverse myelitis (complete or partial), e.g. viral infection, HTLV-1 infection, tabes dorsalis;
- tumour (primary, secondary);
- syringomyelia;
- infarction, e.g. anterior spinal artery syndrome;
- metabolic causes: vitamin $B_{12}$ deficiency producing subacute combined degeneration of the cord.
Imaging of the cord, ideally with MRI, may be helpful in defining the cause of myelopathy.

**References**


**Cross References**

Brown–Séquard syndrome; Lower motor neurone (LMN) syndrome; Paraparesis; Proprioception; Sacral sparing; Suspended sensory loss; Upper motor neurone (UMN) syndrome; Vibration

**Myerson’s Sign**

- see GLABELLAR TAP REFLEX

**Myoclonus**

Myoclonus is involuntary, ‘shock-like’, muscle jerking, arrhythmic more often than regular, of central nervous system (CNS) origin. This may be focal, multifocal, or generalized. Multiple irregular asynchronous myoclonic jerks may be termed polymyoclonus. Myoclonus may be characterized in several ways:

- **Clinical classification (by observation, examination):**
  - Spontaneous
  - *Action or intention:* following voluntary action; may be elicited by asking patient to reach out to touch the examiner’s hand
  - *Reflex, stimulus-sensitive:* jerks produced by somaesthetic stimulation of a limb, in response to loud noises

- **Anatomical/pathophysiological classification (by electrophysiological recordings):**
  - Cortical
  - Subcortical/reticular
  - Propriospinal/segmental

- **Aetiological classification:**
  - *Physiological,* e.g. 'sleep starts' (hypnic jerks)
  - *Essential:* in the absence of any other abnormality of the CNS
  - *Epileptic:* as a manifestation of idiopathic epilepsy
  - *Symptomatic:* of other neurological diseases, of which there are many, including
    - Anoxic brain injury (Lance–Adams syndrome)
    - Cerebrovascular lesions
    - Neoplasia
    - Encephalopathies: especially of metabolic origin (hepatic, renal), but also toxic, viral, paraneoplastic, mitochondrial
    - Degenerations: basal ganglia, spinocerebellar
    - Malabsorption syndromes: coeliac disease, Whipple’s disease
Myoclonus

Storage disorders, e.g. Lafora body disease, Tay–Sachs disease, sialidosis
Dementias: Alzheimer’s disease (usually late), prion disease (usually early in sporadic Creutzfeldt–Jakob disease); not seen in frontotemporal lobar degenerations
Inherited disorders: myoclonus–dystonia syndrome (DYT11)

The clinical differential diagnosis of myoclonus includes chorea, tic, tremor (especially with rhythmic myoclonus), and certain peripheral nerve disorders (fasciculation, myokymia). Periodic limb movement disorder or periodic leg movements of sleep, frequently found in association with restless legs syndrome, is sometimes called ‘nocturnal myoclonus’. Brief lapses of muscle contraction with loss of posture are in some ways the converse of myoclonus and have in the past been labelled ‘negative myoclonus’, although the term asterixis is now preferred.

Drugs useful in the treatment of myoclonus include clonazepam, sodium valproate, primidone, and piracetam. These may need to be given in combination to suppress severe action myoclonus.

References

Cross References
Asterixis; Chorea, Choreoathetosis; Dystonia; Fasciculation; Hiccups; Jactitation; Myokymia; Palatal tremor; Tic; Tremor

Myoedema

Myoedema, or muscle mounding, provoked by mechanical stimuli or stretching of muscle, is a feature of rippling muscle disease, in which the muscle contractions are associated with electrical silence. It has also been reported as a neuromuscular feature of hypothyroidism.

Reference

Myokymia

Myokymia is an involuntary, spontaneous, wave-like, undulating, flickering movement within a muscle (cf. fasciculation); it may be likened to a ‘bag of worms’. Neurophysiologically this corresponds to regular groups of motor unit discharges of peripheral nerve origin. Myokymia is thus related to neuromyotonia and stiffness, since there may be concurrent impairment of muscle relaxation and a complaint of muscle cramps.
A syndrome of superior oblique myokymia is described, often following superior oblique palsy, which produces a microtremor of the eye and causes oscillopsia or transient diplopia. Facial myokymia is a rare facial dyskinesia, possibly related to disinhibition of the facial (VII) nerve nucleus by focal pontine lesions (tumour, demyelination).

Neurophysiological evidence of myokymia may be helpful in the assessment of a brachial plexopathy, since this is found in radiation-induced, but not neoplastic, lesions.

References

Cross References
Fasciculation; Myotonia; Neuromyotonia; Stiffness

Myopathy
The term myopathy means a primary disorder of muscle causing wasting and/or weakness in the absence of sensory abnormalities. Clinically, myopathic processes need to be differentiated from neuropathies, particularly anterior horn cell diseases and motor neuropathies, and neuromuscular junction disorders. Generally in primary muscle disease there are no fasciculations, reflexes are lost late, and phenomena such as (peripheral) fatigue and facilitation do not occur. Myopathies may be subdivided according to the clinical pattern of weakness, and/or their aetiology:

• **Proximal:**
  Affecting shoulder abductors, hip flexors predominantly:
  - Inflammatory: polymyositis, dermatomyositis
  - Progressive muscular dystrophies: Duchenne, Becker, limb-girdle, facioscapulohumeral (FSH)
  - Metabolic: acid maltase deficiency; thyroid dysfunction, Cushing’s syndrome
  - Non-metastatic feature of malignant disease

• **Distal:**
  An unusual pattern for myopathy, which needs to be differentiated from distal polyneuropathy:
  - Myotonic dystrophy
  - Miyoshi dystrophy
  - Desmin myopathy

• **Bulbar palsy**
• **Facial paresis**
• **Diaphragm weakness:**
  - Acid maltase deficiency
  - Acute polymyositis
  - Neuralgic amyotrophy
Myorhythmia

- **Axial myopathy:**
  - Camptocormia (‘bent spine syndrome’)
  - Dropped head syndrome

**References**

**Cross References**
Atrophy; Bulbar palsy; Camptocormia; Dropped head syndrome; Facial paresis; Facial weakness; Fatigue; Gowers’ sign; Paradoxical breathing; Wasting; Weakness

**Myorhythmia**
Myorhythmia is an involuntary movement disorder characterized by rhythmic contraction (1–3 Hz) of muscles producing a coarse tremor, which may affect limbs, face, palate, head, jaw, neck, tongue, eyes, or trunk. The movements are continuous and persist during sleep. They are associated with brainstem or thalamic vascular disease, trauma, alcohol-related nutritional deficiency, phenytoin intoxication, Hashimoto’s encephalopathy, paraneoplasia, and Whipple’s disease.

Although very rare, oculomasticatory myorhythmia is of diagnostic importance since it is pathognomonic for Whipple’s disease of the nervous system. Characteristically there is also convergent–divergent pendular nystagmus with synchronous rhythmic movement of the mouth, tongue, jaw, and sometimes proximal and distal skeletal muscles. The neurological manifestations of Whipple’s disease are protean, and include dementia, ataxia, supranuclear ophthalmoplegia (with sparing of the pupils), epileptic seizures, myoclonus, nystagmus, and psychosis. The condition is caused by the bacterium *Tropheryma whippelii*. Treatment is with antibiotics, usually a 2-week intravenous course of trimethoprim–sulphamethoxazole or ceftriaxone followed by oral treatment for 1 year. Sodium valproate may be helpful for the involuntary movements which do not respond to antibiotics.

**References**

**Cross References**
Ataxia; Dementia; Myoclonus; Nystagmus

**Myotonia**
Myotonia is a stiffness of muscles with inability to relax after voluntary contraction (action myotonia), or induced by electrical or mechanical (e.g. percussion
myotonia) excitation. The phenomenon is often described by patients as ‘cramp’ or stiffness. This is a reflection of primary muscle disease (i.e. myogenic; cf. neuromyotonia, neurogenic muscle stiffness, peripheral nerve hyperexcitability), which persists after peripheral nerve or neuromuscular junction blockade.

Neurophysiology reveals myotonic discharges, with prolonged twitch relaxation phase, which may be provoked by movement, percussion, and electrical stimulation of muscle; discharges typically wax and wane.

Myotonia may be aggravated by hyperkalaemia, depolarizing neuromuscular blocking drugs (e.g. suxamethonium), and anticholinesterase drugs (neostigmine). Other factors that can induce myotonia include hypothermia, mechanical or electrical stimulation (including surgical incision and electrocautery), shivering, and use of inhalational anaesthetics.

A similar clinical phenomenon of slow muscle relaxation may be observed in other circumstances, for example hypothyroidism, but without the characteristic EMG findings of myotonia, hence this is labelled as pseudomyotonia. Paramyotonia is myotonia exacerbated by cold and exertion (paradoxical myotonia).

Recognized causes of myotonia include

- myotonic dystrophy types 1 and 2;
- hyperkalaemic periodic paralysis;
- myotonia congenita (autosomal dominant Thomsen’s disease, autosomal recessive Becker’s myotonia);
- K⁺-aggravated myotonia;
- Schwartz–Jampel syndrome (chondrodystrophic myotonia).

Mutations in genes encoding voltage-gated ion channels have been identified in some of the inherited myotonias, hence these are channelopathies: skeletal muscle voltage-gated Na⁺ channel mutations have been found in K⁺-aggravated myotonia, and also paramyotonia congenita and hyperkalaemic periodic paralysis. Chloride (Cl⁻) channel mutations have been identified in myotonia congenita. These latter conditions respond best to mexiletine.

References

Cross References
Neuromyotonia; Paramyotonia; Percussion myotonia; Pseudomyotonia; Stiffness; Warm-up phenomenon; Woltman’s sign

- 235 -
Narcolepsy, Narcoleptic Syndrome  
- see HYPERSOMNOLENCE

Nasopalpebral Reflex  
- see GLABELLAR TAP REFLEX

Negative Myoclonus  
- see ASTERIXIS

Negative Tremor  
- see ASTERIXIS

Negativism  
Negativism is a motor sign of mental disorder, usually schizophrenia, consisting of the patient doing the opposite of what is asked and actively resisting efforts to persuade compliance. Movement of a limb in response to application of pressure despite the patient having been told to resist (mitgehen) is one element of negativism. It may also be a feature of catatonia. The similarity of some of these features to gegenhalten suggests the possibility of frontal lobe dysfunction as the underlying cause.

Cross References  
Catatonia; Gegenhalten

Neglect  
Neglect is a failure to orient towards, respond to, or report novel or meaningful stimuli. If failure to respond can be attributed to concurrent sensory or motor deficits (e.g. hemiparesis, hemianopia, visuospatial deficits) neglect is not present.

Neglect can involve stimuli in the extrapersonal environment (e.g. visual neglect) or personal space (e.g. personal neglect or asomatognosia). This dichotomy may also be characterized as egocentric (neglecting hemispace defined by the midplane of the body) and allocentric (neglecting one side of individual stimuli). Neglect of contralateral hemispace may also be called unilateral spatial neglect, hemi-inattention, or hemineglect. Lesser degrees of neglect may be manifest as extinction (double simultaneous stimulation). Motor neglect may be evident as hemiakinesia, hypokinesia, or motor impersistence. Alexia may sometimes be a consequence of neglect (neglect alexia). Alloaesthesia and allokinesia may also be features of neglect.

Neglect may be obvious (e.g. patient not dressing one side of the body), but is sometimes more subtle, in which case it may be tested for using various simple tests:
Neglect is more common after right rather than left brain damage, usually of vascular origin. The angular gyrus and parahippocampal gyrus may be central to the development of visual neglect. Marked degrees of neglect may seriously hamper attempts at neurorehabilitation.

References

Cross References
Alexia; Alloaesthesia; Allokinesia; Asomatognosia; Eastchester clapping test; Extinction; Hemiakinesia; Hypokinesia; Impersistance

Negro’s Sign
Negro has two eponymous signs:

- Cogwheel (jerky) type of rigidity in basal ganglia disorders;
- In both peripheral and central facial paralyses, the eyeball deviates outwards and elevates more than normal when the patient attempts to look up due to overaction of the inferior oblique and superior rectus muscles, respectively.

Reference

Cross References
Bell’s palsy; Facial paresis; Parkinsonism; Rigidity

Neologism
A neologism is a non-word approximating to a real word, produced in spontaneous speech; it is thought to result from an inability to organize phonemes appropriately in the process of speech production. Hence, this is a type of literal or phonemic paraphasia encountered in aphasic syndromes, most usually those resulting from left superior temporal lobe damage (Wernicke type). (The word ‘scientist’ is said to be a neologism coined in the nineteenth century by William Whewell.)

Cross References
Aphasia; Paraphasia; Schizophrenia; Wernicke’s aphasia
Neuromyotonia

Neri's Test
- see LASÈGUE’S SIGN

Nerve Thickening
The characterization of a peripheral neuropathy should always include examination to see if any nerves are thickened. Good places to feel for nerve thickening include the elbow (ulnar nerve), anatomical snuffbox (superficial radial nerves), and head of the fibula (common peroneal nerve). Nerve thickening may be noted in a variety of conditions, in some by examination, in others using imaging techniques:

- Leprosy
- Hereditary motor and sensory neuropathies (HMSN), especially types I, III, and IV (Refsum’s disease)
- Hereditary neuropathy with liability to pressure palsies (HNLPP)/tomaculous neuropathy
- Neurofibromatosis 1
- Sarcoidosis
- Chronic inflammatory demyelinating neuropathy/ophthalmoplegic migraine
- Nerve tumours (localized)
- Amyloidosis (familial amyloid polyneuropathy, primary systemic amyloidosis): rare

References
Donaghy M. Enlarged peripheral nerves. Practical Neurology 2003; 3: 40–45.

Cross Reference
Neuropathy

Neuromyotonia
Neuromyotonia is neurogenic muscle stiffness (cf. myotonia, myogenic muscle stiffness) which reflects peripheral nerve hyperexcitability. Clinically this is manifest as muscle cramps and stiffness, particularly during and after muscle contraction, and as muscular activity at rest (myokymia, fasciculations). Tendon areflexia and abnormal postures of hands and feet may also be observed. Sensory features such as paraesthesias and central nervous system features (Morvan’s syndrome) can occur. A syndrome of ocular neuromyotonia has been described in which spasms of the extraocular muscles cause a transient heterophoria and diplopia.

Physiologically neuromyotonia is characterized by continuous motor unit and muscle fibre activity which is due to peripheral nerve hyperexcitability; it is abolished by curare (cf. myotonia). Spontaneous firing of single motor units as doublet, triplet, or multiplet discharges with high-intraburst frequency (40–300/s) at irregular intervals is the hallmark finding.

Neuromyotonia may be associated with autoantibodies directed against presynaptic voltage-gated K⁺ channels. Around 20% of patients have an
Neuronopathy underling small cell lung cancer or thymoma, suggesting a paraneoplastic aetiology in these patients. Neuromyotonia has also been associated with mutations within the voltage-gated K\(^+\) ion channel gene.

Neuromyotonia usually improves with symptomatic treatments such as carbamazepine, phenytoin, lamotrigine, and sodium valproate, in combination if necessary. Paraneoplastic neuromyotonia often improves and may remit after treatment of the underlying tumour.

References

Cross References
Fasciculation; Myokymia; Myotonia; Paramyotonia; Pseudomyotonia; Stiffness

Neuroneopathy
Neuroneopathies are disorders affecting neuronal cell bodies in the ventral (anterior) horns of the spinal cord or dorsal root ganglia, hence motor and sensory neuronopathies, respectively. Sensory neuronopathies (ganglionopathy, polyganglionopathy) have a more limited differential diagnosis than neuropathies, including

- Paraneoplasia: anti-Hu antibody syndrome (although a similar syndrome, presumed paraneoplastic, may occur in the absence of these antibodies)
- Sjögren's syndrome
- Associated with anti-GD1b ganglioside antibodies
- CIDP
- HIV

Anterior horn cell (motor neurone) disorders may be classified as motor neuronopathies, including motor neurone disease (amyotrophic lateral sclerosis), spinal muscular atrophies, and poliomyelitis.

Cross Reference
Neuropathy

Neuropathy
Neuropathies are disorders of peripheral nerves. Various clinical patterns of peripheral nerve involvement may be seen:

- *Mononeuropathy*: sensory and/or motor involvement in the distribution of a single nerve.
- *Mononeuropathy multiplex*: simultaneous involvement of two or more nerves, usually in different parts of the body; if due to inflammatory disease, as is often the case, this may be described as mononeuritis multiplex.
Polyneuropathy: a widespread process, predominantly affecting the distal parts of nerves; may be predominantly sensory ('glove and stocking' sensory loss) or motor, with or without concomitant autonomic involvement.

These clinical patterns may need to be differentiated in practice from disorders affecting the neuronal cell bodies in the ventral (anterior) horns of the spinal cord or dorsal root ganglia (motor and sensory neuronopathies, respectively); and disorders of the nerve roots (radiculopathy) and plexuses (plexopathy). Clinical signs resulting from neuropathies are of lower motor neurone type (wasting, weakness, reflex diminution, or loss).

The causes of neuropathy are many. Mononeuropathies often result from local compression (entrapment neuropathy), trauma, or diabetes. Mononeuropathy multiplex often reflects intrinsic inflammation (e.g. polyarteritis nodosa, Churg–Strauss syndrome, systemic lupus erythematosus, rheumatoid arthritis, Sjögren’s syndrome, cryoglobulinaemia, isolated PNS vasculitis). Polyneuropathies may have genetic, infective, inflammatory, toxic, nutritional, and endocrine aetiologies. Many neuropathies, particularly polyneuropathies in the elderly, remain idiopathic or cryptogenic, despite intensive investigation.

References

Cross References
Amyotrophy; Lower motor neurone (LMN) syndrome; Neuronopathy; Plexopathy; Radiculopathy; Wasting; Weakness

Nominal Aphasia
- see ANOMIA

Nuchal Rigidity
Nuchal rigidity is neck stiffness and is usually synonymous with meningism, in which case other signs of meningeal irritation are usually present (Kernig's sign, Brudzinski's neck sign). If these other signs are absent, then isolated nuchal rigidity may suggest a foraminal pressure cone. It may also occur in syndromes causing predominantly axial (as opposed to limb) rigidity (e.g. progressive supranuclear palsy). In intubated patients, there may be resistance to passive neck movements.

Cross References
Brudzinski's (neck) sign; Kernig’s sign; Meningism; Parkinsonism

Nuchocephalic Reflex
In a standing subject, rapid turning of the shoulders to either left or right (eyes closed to avoid fixation) is associated with bilateral contraction of the cervical musculature so that the head is held in the original position. This nuchocephalic reflex...
Nyctalopia

Nyctalopia, or night blindness, is an impairment of visual acuity specific to scotopic vision, implying a loss or impairment of rod photoreceptor function. Patients may spontaneously complain of a disparity between daytime and nocturnal vision, in which case acuity should be measured in different ambient illumination. Nyctalopia may be a feature of:

- Retinitis pigmentosa
- Vitamin A deficiency
- Cancer-associated retinopathy: most commonly associated with small cell lung cancer (antirecoverin antibodies may be detected), though gynaecological malignancy and melanoma have also been associated (with antibipolar retinal cell antibodies in the latter).

Cross References
Hemeralopia; Retinitis pigmentosa

Nylen–Bárány Manoeuvre
- see HALLPIKE MANOEUVRE, HALLPIKE TEST

Nystagmoid Jerks
- see NYSTAGMUS

Nystagmus
Nystagmus, or talantropia, is an involuntary to-and-fro oscillatory movement of the eyeballs, of which there are many varieties. It is usually bilateral, but occasionally may be unilateral, as in internuclear ophthalmoplegia (INO). The pathophysiological underpinnings are diverse, but all involve brainstem nuclei and tracts which control eye movements and gaze holding, especially the III, IV, and VI cranial nerve nuclei, paramedian pontine reticular formation, vestibular nuclei, medial longitudinal fasciculus, central tegmental tract, cerebellar connections to these structures, interstitial nucleus of Cajal, and nucleus prepositus hypoglossi. The nature of the nystagmus may permit inferences about the precise location of pathology. Observations should be made in the nine cardinal positions of gaze for direction, amplitude, and beat frequency of nystagmus. Nystagmus may be abortive or sustained in duration.
Nystagmus may be classified in various ways:

- **Physiological:**
  - Optokinetic nystagmus (OKN; e.g. looking out of a moving railway carriage)
  - Induced by vestibular stimuli (e.g. merry-go-round; caloric testing)
  - Nystagmoid jerks: in extremes of lateral or vertical gaze (end-point nystagmus, a form of gaze-evoked nystagmus)

- **Pathological:**
  - Nystagmus may be classified according to direction, waveform, anatomy/aetiology, or clinical frequency (common, rare)

It is important to distinguish nystagmus from other involuntary eye movements such as square wave jerks, ocular flutter, and opsoclonus.

- **Directional classification of nystagmus:**
  - Horizontal (common)
  - Vertical (rare):
    - **Downbeat:** seen not only with structural lesions of the cervico-medullary junction, midline cerebellum, and floor of the fourth ventricle, but also with more diffuse cerebellar disease
    - **Upbeat:** of less localizing value than downbeat; upbeat nystagmus may occur with pontomesencephalic, pontomedullary, and even caudal medullary lesions (infarct, inflammation); bow-tie nystagmus is probably a variant of upbeat nystagmus
    - **Torsional:** usually accompanies horizontal nystagmus of peripheral vestibular (labyrinthine) origin

- **Waveform classification of nystagmus:**
  - **Jerk nystagmus:**
    - At least one of the directions of eye movement is slow (slow phase; \(<40^\circ/s\)) followed by a rapid, corrective, saccadic movement in the opposite direction (fast phase) for which direction the nystagmus is named.
    - However, since it is the slow phase which is pathological, it is more eloquent concerning anatomical substrate. The intensity of jerk nystagmus may be classified by a scale of three degrees:
      - 1st degree: present when looking in the direction of the fast phase;
      - 2nd degree: present in the neutral position;
      - 3rd degree: present when looking in the direction of the slow phase (i.e. present in all directions of gaze).
  - **Pendular or undulatory nystagmus:**
    - In which the movements of the eyes are more or less equal in amplitude and velocity (sinusoidal oscillations) about a central (null) point.
      - This is often congenital, may be conjugate or disconjugate (sometimes monocular), but is not related to concurrent internuclear ophthalmoplegia or asymmetry of visual acuity.
      - Acquired causes include multiple sclerosis and brainstem infarctions.

When studied using oculography, the slow phase of jerk nystagmus may show a uniform velocity (‘saw-toothed’), indicative of imbalance in vestibulo-ocular...
Nystagmus

reflex activity. A slow phase with exponentially decreasing velocity (negative exponential slow phase) is ascribed to ‘leakiness’ of a hypothetical neural integrator, a structure which converts eye or head velocity signals into approximations of eye or head position signals (thought to lie in the interstitial nucleus of Cajal in the midbrain for vertical eye movements and in the nucleus propositus hypoglossi for horizontal eye movements). A slow phase with exponentially increasing velocity (high-gain instability, runaway movements) may be seen in congenital or acquired pendular nystagmus. The pathophysiology of acquired pendular nystagmus is thought to be deafferentation of the inferior olive by lesions of the red nucleus, central tegmental tract, or medial vestibular nucleus.

- **Anatomical/aetiological classification of nystagmus:**

  - **Peripheral vestibular:**
    
    unidirectional (directed to side opposite lesion), and more pronounced when looking in direction of the fast phase (i.e. 1st degree), usually with a rotatory component and associated with vertigo. Tends to fatigue, and usually transient (e.g. in Hallpike manoeuvre). Nystagmus of peripheral vestibular origin is typically reduced by fixation (hence these patients hold their heads still) and enhanced by removal of visual fixation (in the dark, with Frenzel’s lenses).

  - **Central vestibular:**
    
    unidirectional or multidirectional, 1st, 2nd or 3rd degree; typically sustained and persistent. There may be other signs of central pathology (e.g. cerebellar signs, upper motor neurone signs). Not affected by removal of visual fixation.

  - **Cerebellar/brainstem:**
    
    commonly gaze-evoked due to a failure of gaze-holding mechanisms. It may be unidirectional with a unilateral cerebellar lesion (e.g. vascular disease) in which case it typically occurs when the eyes are looking in the direction of the lesion (cf. peripheral vestibular nystagmus); multidirectional nystagmus of cerebellar origin may occur in multiple sclerosis, drug/toxin exposure, cerebellar degenerations.

  - **Congenital:**
    
    usually horizontal, pendular-type nystagmus; worse with fixation, attention, and anxiety. It may appear with blindness of childhood onset or be acquired with neurological disease (multiple sclerosis, mitochondrial disease, Whipple’s disease, Pelizaeus–Merzbacher disease).

Other forms of nystagmus include

- **Ataxic/dissociated:**
  
  in abducting >> adducting eye, as in internuclear ophthalmoplegia and pseudointernuclear ophthalmoplegia.

- **Periodic alternating:**
  
  primary position nystagmus, almost always in the horizontal plane, which stops and then reverses direction every minute or so; 4–5 min observation may be required to see the whole cycle; localizing value similar to downbeat nystagmus.
Nystagmus

- **Convergence–retraction (Körber–Salus–Elschnig syndrome):**
  
  adducting saccades (medial rectus contraction), occurring spontaneously or on attempted upgaze, often accompanied by retraction of the eyes into the orbits, associated with mesencephalic lesions of the pretectal region (e.g. pinealoma).

- **See-saw:**
  
  a disconjugate cyclic movement of the eyes, comprising elevation and intorsion of one eye while the other eye falls and extorts, followed by reversal of these movements; may be congenital (e.g. with albinism, retinitis pigmentosa) or acquired (mesodiencephalic or lateral medullary lesions, e.g. brainstem stroke, head trauma, syringobulbia).

Many pathologies may cause nystagmus, the most common being demyelination, vascular disease, tumour, neurodegenerative disorders of cerebellum and/or brainstem, metabolic causes (e.g. Wernicke–Korsakoff’s syndrome), paraneoplasia, drugs (alcohol, phenytoin, barbiturates, sedative-hypnotic drugs), toxins, and epilepsy.

Treatment of nystagmus is usually that of the underlying cause, where possible. Pendular nystagmus may respond to anticholinesterases, consistent with its being a result of cholinergic dysfunction. Periodic alternating nystagmus responds to baclofen, hence the importance of making this diagnosis. See-saw nystagmus may respond to baclofen, clonazepam, or alcohol.

**References**


**Cross References**

Caloric testing; Hallpike manoeuvre, Hallpike test; Internuclear ophthalmoplegia (INO); Myorhythmia; Optokinetic nystagmus (OKN), Optokinetic response; Opsoclonus; Oscillopsia; Palatal myoclonus; Pendular nystagmus; Pseudointernuclear ophthalmoplegia; Spasmus nutans; Square wave jerks; Vertigo
Obscurations
Visual obscurations are transient losses (‘greying out’) of vision lasting a few seconds, occurring in the context of raised intracranial pressure (ICP), and especially associated with activities known to elevate ICP, such as coughing, sneezing, bending down, straining at stool, and relieved by their cessation. These symptoms are thought to reflect critical compromise of optic nerve head perfusion and are invariably associated with the finding of papilloedema. Obscurations mandate urgent investigation and treatment to prevent permanent visual loss. Transient visual obscurations may occasionally be due to optic disc drusen.

Cross Reference
Papilloedema

Obtundation
Obtundation is a state of altered consciousness characterized by reduced alertness and a lessened interest in the environment, sometimes described as psychomotor retardation or torpor. An increased proportion of time is spent asleep and the patient is drowsy when awake. Obtundation is a less severe impairment of consciousness than stupor.

Cross References
Coma; Psychomotor retardation; Stupor

Ocular Apraxia
Ocular apraxia (ocular motor apraxia) is a disorder of voluntary saccade initiation; reflexive saccades and spontaneous eye movements are preserved. Ocular apraxia may be overcome by using dynamic head thrusting, with or without blinking (to suppress vestibulo-ocular reflexes): the desired fixation point is achieved through reflex contraversive tonic eye movements to the midposition following the overshoot of the eyes caused by the head thrust.

The anatomical substrate of ocular apraxia is not certain. Ocular apraxia may occur as a congenital syndrome (in the horizontal plane only: Cogan’s syndrome), or may be acquired in ataxia telangiectasia (Louis–Bar syndrome), Niemann–Pick disease (mainly vertical plane affected), and Gaucher’s disease (horizontal plane only).

Cross References
Apraxia; Saccades

Ocular Bobbing
Ocular bobbing refers to intermittent abnormal vertical eye movements, usually conjugate, consisting of a fast downward movement followed by a slow return to the initial horizontal eye position. The sign has no precise localizing value, but is most commonly associated with intrinsic pontine lesions, e.g. infarct,
haemorrhage, tumour, central pontine myelinolysis. It has also been described in encephalitis, Creutzfeldt–Jakob disease, and toxic encephalopathies. Its pathophysiology is uncertain but may involve mesencephalic and medullary burst neurone centres. Variations on the theme include

- **Inverse ocular bobbing**: slow downward movement, fast return (also known as fast upward ocular bobbing or ocular dipping);
- **Reverse ocular bobbing**: fast upward movement, slow return to midposition;
- **Converse ocular bobbing**: slow upward movement, fast down (also known as slow upward ocular bobbing or reverse ocular dipping).

**References**

**Cross Reference**
Ocular dipping

**Ocular Dipping**
Ocular dipping, or inverse ocular bobbing, consists of a slow spontaneous downward eye movement with a fast return to the midposition. This may be observed in anoxic coma or following prolonged status epilepticus and is thought to be a marker of diffuse, rather than focal, brain damage. Reverse ocular dipping (slow upward ocular bobbing) consists of a slow upward movement followed by a fast return to the midposition.

**Reference**

**Cross Reference**
Ocular bobbing

**Ocular Flutter**
Ocular flutter is an eye movement disorder characterized by involuntary bursts of back-to-back horizontal saccades without an intersaccadic interval (cf. square wave jerks). Ocular flutter may be accurately diagnosed with oculography. The postulated mechanism of ocular flutter is loss of ‘pause’ neuronal inhibition of ‘burst’ neurone function in the paramedian pontine reticular formation (PPRF). A case of ocular flutter with a circumscribed inflammatory pontine lesion involving the PPRF, in which clinical and neuroradiological improvement occurred together, has been reported, supporting the argument that, at least in some cases, PPRF lesions may be associated with ocular flutter.

**Reference**

**Cross References**
Opsoclonus; Saccades; Saccadic intrusion, Saccadic pursuit; Square wave jerks

**Ocular Myoclonus**
- see MYOCLONUS; PALATAL MYOCLONUS
Oculogyric Crisis

Ocular Tilt Reaction
The ocular tilt reaction is a postural synkinesis consisting of the triad of:

• ocular torsion;
• lateral head tilt to the same side;
• skew deviation with hypotropia ipsilateral to the direction of head/eye torsion.

The ocular tilt reaction (OTR) is due to disordered function of one utricle or its brainstem connections (vestibular nerve, vestibular nuclei, medial longitudinal fasciculus, interstitial nucleus of Cajal), hence a brainstem otolith-ocular reflex. It has occasionally been reported with cerebellar lesions and may be under inhibitory cerebellar control. OTR may be tonic, as in the lateral medullary syndrome, or paroxysmal, as in multiple sclerosis.

Reference

Cross References
Hypotropia; Lateral medullary syndrome; Skew deviation; Synkinesia, Synkinesis; Tullio phenomenon; Vestibulo-ocular reflexes

Oculocephalic Response
Oculocephalic responses are most commonly elicited in unconscious patients; the head is passively rotated in the horizontal or vertical plane (doll’s head manoeuvre) and the eye movements are observed. Conjugate eye movement in a direction opposite to that in which the head is turned is indicative of an intact brainstem (intact vestibulo-ocular reflexes). With pontine lesions, the oculocephalic responses may be lost, after roving eye movements but before caloric responses disappear.

Cross References
Caloric testing; Coma; Doll’s head manoeuvre, Doll’s eye manoeuvre; Head impulse test; Roving eye movements; Supranuclear gaze palsy; Vestibulo-ocular reflexes

Oculogyric Crisis
Oculogyric crisis is an acute dystonia of the ocular muscles, usually causing upward and lateral displacement of the eye. It is often accompanied by a disorder of attention (obsessive, persistent thoughts), with or without dystonic or dyskinetic movements. It occurs particularly with symptomatic (secondary), as opposed to idiopathic (primary), dystonias, for example, postencephalitic and neuroleptic-induced dystonia, the latter now being the most common cause. This is usually an acute effect but may on occasion be seen as a consequence of chronic therapy (tardive oculogyric crisis). It has also been described with Wilson’s disease, neuroleptic malignant syndrome, and organophosphate poisoning. Lesions within the lentiform nuclei have been recorded in cases with oculogyric crisis. Treatment of acute neuroleptic-induced dystonia is either parenteral benzodiazepine or an anticholinergic agent such as procyclidine, benztropine, or trihexyphenidyl.
Oculomasticatory Myorhythmia

References

Cross References
Dyskinesia; Dystonia

Oculomasticatory Myorhythmia
- see MYORHYTHMIA

Oculomotor (III) Nerve Palsy
Oculomotor (III) nerve palsy produces
- **Ptosis:** weakness of levator palpebrae superioris (LPS), +/- Müller’s muscle;
- **Mydriasis:** impaired parasympathetic outflow to the pupil (‘internal ophthalmoplegia’); most obvious in a well-lit room (cf. Horner’s syndrome);
- **Diplopia:** weakness of medial rectus (MR), inferior rectus (IR), superior rectus (SR), and inferior oblique (IO) muscles causing the eye to point ‘down and out’ (external ophthalmoplegia); the presence of intorsion confirms integrity of superior oblique muscle/trochlear (IV) nerve function.

These changes may be complete or partial.

Pathological correlates of third nerve palsy may occur anywhere from the brainstem to the orbit:
- **Intramedullary (brainstem):**
  - **Nuclear:** very rare; SR subnucleus lesion causes bilateral denervation; other clinical signs may be expected, such as pupillary (Edinger–Westphal nucleus) and medial longitudinal fasciculus involvement.
  - **Fascicular (within substance of midbrain):** all muscles or specific muscles involved, + other clinical signs expected, such as contralateral ataxia (Claude’s syndrome), hemiparesis (Weber’s syndrome).
- **Extramedullary:**
  - **Subarachnoid space:** peripherally located pupillomotor fibres often spared by ischaemic lesions, but not by space-occupying lesions (e.g. aneurysm), however, the distinction is not absolute.
  - **Cavernous sinus:** III runs over trochlear nerve; other oculomotor nerves +/- trigeminal nerve often affected.
  - **Superior orbital fissure:** superior division/ramus to SR, LPS; inferior to MR, IR, IO; selective involvement (divisional palsy) may occur; proptosis with space-occupying lesions.
  - **Orbit:** paresis of isolated muscle almost always from orbital lesion or muscle disease.

Oculomotor nerve palsies may be distinguished as ‘pupil involving’ or ‘pupil sparing’, the former implying a ‘surgical’, the latter a ‘medical’ cause, but this distinction only holds for complete palsies. Incomplete palsies are more likely to be of ‘surgical’ origin (e.g. posterior communicating artery aneurysm). Neuroimaging is the appropriate management if in doubt. Transtentorial (uncal)
herniation due to raised intracranial pressure may, particularly in its early stages, cause an oculomotor nerve palsy due to stretching of the nerve, a ‘false-localizing sign’.

References

Cross References
Diplopia; Divisional palsy; ‘False-localizing signs’; Hutchinson’s pupil; Mydriasis; Ophthalmoparesis, Ophthalmoplegia; Ptosis; Pupil sparing

Oculovestibular Response
- see CALORIC TESTING; VESTIBULO-OCULAR REFLEXES

Okay Sign
- see ‘PINCH SIGN’

One-and-a-Half Syndrome
The one-and-a-half syndrome consists of an ipsilateral horizontal gaze palsy and an ipsilateral internuclear ophthalmoplegia, such that the only preserved horizontal eye movement is abduction in one eye; vertical movements and convergence are spared. This results from a brainstem lesion which involves both the abducens (VI) nerve nucleus or paramedian pontine reticular formation, causing ipsilateral horizontal gaze palsy, and the adjacent medial longitudinal fasciculus, causing internuclear ophthalmoplegia. In young patients this is most often due to demyelination, in the elderly to brainstem ischaemia; brainstem arteriovenous malformation or tumour may also be responsible. Myasthenia gravis may cause a pseudo-one-and-a-half syndrome.

A vertical one-and-a-half syndrome has also been described, characterized by vertical upgaze palsy and monocular paresis of downgaze, either ipsilateral or contralateral to the lesion.

References

Cross References
Eight-and-a-half syndrome; Gaze palsy; Internuclear ophthalmoplegia (INO)

Onion Peel, Onion Skin
These terms have been used to describe the pattern of facial sensory loss with perioral sparing (Dejerine pattern), seen with intramedullary or cervicomедullary lesions and with tabes dorsalis. It reflects the somatotopic sensory representation in the spinal nucleus of the trigeminal nerve: midline face (nose, mouth) represented rostrally, lateral facial sensation represented caudally. The pattern of sensory impairment has also been termed ‘balaclava helmet’.
Ophthalmoparesis, Ophthalmoplegia

Ophthalmoparesis is a weakness or limitation, ophthalmoplegia a paralysis, of eye movements. Causes may be central (CNS pathways), or peripheral (cranial nerve nuclei, cranial nerves, neuromuscular junction, extraocular muscles). A distinction is sometimes drawn between:

- **External ophthalmoplegia:**
  
  weakness of the extraocular muscles of central, neuromuscular, or myopathic origin:
  
  - *Supranuclear*: e.g. progressive supranuclear palsy, abetalipoproteinemia;
  - *Nuclear, internuclear*: e.g. internuclear ophthalmoplegia (INO), Möbius syndrome;
  - *Cranial nerve palsy*: III, IV, VI, or combinations thereof;
  - *Neuromuscular junction*: myasthenia gravis;
  - *Extraocular muscles*: e.g. oculopharyngeal muscular dystrophy (OPMD), chronic progressive external ophthalmoplegia (CPEO), thyroid ophthalmopathy.

  The term ‘ophthalmoplegia plus’ has been used to denote the combination of progressive external ophthalmoplegia with additional symptoms and signs, indicative of brainstem, pyramidal, endocrine, cardiac, muscular, hypothalamic, or auditory system involvement, as in mitochondrial disease.

- **Internal ophthalmoplegia:**
  
  fixity of the pupil with loss of all pupillary reflexes (iridoplegia) and ciliary apparatus.

  Hence in an oculomotor (III) nerve palsy there may be both internal and external ophthalmoplegia.

  If structural disease and myasthenia gravis are excluded, then mitochondrial disorder (CPEO) may be responsible for ophthalmoplegia, even if this is not evident on quadriceps muscle biopsy.

Reference


Cross References

Diplopia; Internuclear ophthalmoplegia (INO); Miosis; Mydriasis; Oculomotor (III) nerve palsy; Pupillary reflexes; Pupil sparing

Opisthotonos

Opisthotonos is an abnormal posture consisting of arching of the back and extension of the limbs such that the body may be supported just on the head and ankles (*arc de cercle*). Opisthotonos may be seen in:

- Coma; decerebrate rigidity
- Basilar meningitis

Cross Reference

Balaclava helmet
Opsoclonus

- Hydrocephalus
- Structural lesions of the posterior fossa
- Cerebellar fits due to intermittent tonsillar herniation
- Acute drug-(neuroleptic-) induced dystonic reaction; or chronic feature of tardive dystonia
- Tetanus
- Syncope (especially in children)
- Metabolic disorders: kernicterus, Gaucher's disease (type II)
- Drug-induced: propofol
- Pseudoseizures

As in decerebrate rigidity, opisthotonos may reflect unopposed extensor tone from the intact vestibular nuclei released from supratentorial control.

Cross References
Coma; Decerebrate rigidity; Emposthotonos

Oppenheim’s Sign
Oppenheim’s sign is a variant method for eliciting the plantar response, by application of heavy pressure to the anterior surface of the tibia, for example, with the thumb, and moving it down from the patella to the ankle. Extension of the halluc (upgoing plantar response, Babinski’s sign) is pathological. Like Chaddock’s sign, Oppenheim’s sign always postdates the development of Babinski’s sign as a reliable indicator of corticospinal pathway (upper motor neurone) pathology.

References

Cross References
Babinski’s sign (1); Chaddock’s sign; Gordon’s sign; Plantar response; Upper motor neurone (UMN) syndrome

Oppenheim’s Useless Hand Sign
- see USELESS HAND OF OPPENHEIM

Opsoclonus
Opsoclonus, or saccadomania, is an eye movement disorder characterized by involuntary bursts of polydirectional saccades (sometimes with a horizontal preference) without an intersaccadic interval (cf. square wave jerks). Like ocular flutter, opsoclonus may be accurately characterized with oculography.

Although some normal individuals can voluntarily induce opsoclonus, generally it reflects mesencephalic or cerebellar disease affecting the omnipause cells which exert tonic inhibition of the burst neurones which generate saccades. Recognized causes of opsoclonus include

- Paraneoplasia: in children with neuroblastoma (Kinsbourne’s syndrome); in adults the opsoclonus–myoclonus syndrome (‘dancing eyes, dancing feet’) is most commonly associated with small cell lung cancer but it may also occur in association with breast cancer in which case onconeural antibodies (anti-Ri, or type 2 antineuronal nuclear antibodies [ANNA-2]) may be detected in serum and CSF.
Postinfectious: a monophasic disorder following respiratory or gastrointestinal infection.

Intraparenchymal (especially mesencephalic) lesions, e.g. tumour, demyelination, neurosarcoidosis, metabolic/toxic encephalopathy.

Postinfectious opsinclonus generally remits spontaneously. Of the paraneoplastic disorders, opsinclonus associated with lung and breast tumours persists and the patients decline from their underlying illness; neuroblastoma associated opsinclonus may be steroid responsive. IVIg, clonazepam, and valproate have also been used as symptomatic treatments.

References

Cross References
Ocular flutter; Saccadic intrusion, Saccadic pursuit; Square wave jerks

**Optic Aphasia**

Optic aphasia is a visual modality-specific naming disorder. It has sometimes been grouped with associative visual agnosia, but these patients are not agnosic since they can demonstrate recognition of visually presented stimuli by means other than naming, e.g. gesture. Moreover, these patients are not handicapped by their deficit in everyday life, whereas agnosic patients are often functionally blind. Objects that are semantically related can be appropriately sorted, indicating intact semantics. This is not simply anomia, since the deficit is specific to visual stimuli; objects presented in tactile modality, or by sound, or by spoken definition, can be named. Naming errors are often semantic, and perseverations (‘*conduit d’approche*’) are common. Perception is intact, evidenced by the ability to draw accurately objects which cannot be named. Reading is poorly performed.

Optic aphasia is associated with unilateral lesions of the left occipital cortex and subjacent white matter.

The neuropsychological explanation of optic aphasia is unclear. It may be a mild type of associative visual agnosia, despite the differences.

References
Beauvois MF. Optic aphasia: a process of interaction between vision and language. *Philosophical Transactions of the Royal Society, Series B* 1982; 298: 35–47.

Cross References
Anomia; *Conduit d’approche*; Visual agnosia
**Optic Ataxia**

Optic ataxia is impaired voluntary reaching for a visually presented target, with misdirection and dysmetria. It may resemble cerebellar ataxia. Visual fixation is possible but reaching under visual guidance is impaired. Tactile search with the palm and fingers may be undertaken in searching for an object, using somatosensory cues to compensate for impaired access to visual information. Hence this may be characterized as a modality-specific apraxia, wherein visual information cannot be used to guide goal-directed movements. The disorder is both retinotopic and somatotopic.

Optic ataxia occurs with lesions of the intraparietal sulcus and regions medial and superior to it; the primary visual cortex is intact. It is one feature, along with psychic paralysis of gaze (‘sticky fixation’) and simultanagnosia (visual disorientation), of Balint’s syndrome in which there is some evidence for parieto-occipital (and possibly frontal) lobe dysfunction (disconnection).

**Reference**


**Cross References**

Apraxia; Ataxia; Balint’s syndrome; Dysmetria; Simultanagnosia; Visual disorientation; Visual form agnosia

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**Optic Atrophy**

Optic atrophy is pallor of the optic nerve head as visualized by ophthalmoscopy. The temporal disc may appear pale in a normal fundus, so that optic atrophy can only be confidently diagnosed when there is also nasal pallor, although temporal pallor may follow damage to the macular fibre bundle with central visual defects.

Optic atrophy may be the consequence of any optic neuropathy which causes optic nerve damage leading to gliotic change of the optic nerve head. Although most often seen with optic nerve pathology, it may be a consequence of pathology in the retina, optic chiasm, or optic tract. ‘Hemianopic’ optic atrophy indicates involvement of the optic tract or lateral geniculate body.

The appearance of optic atrophy is non-specific with respect to aetiology. Recognized causes include:

- Previous optic neuritis
- Chronic papilloedema
- Chronic optic nerve compression (see Foster Kennedy syndrome)
- Hereditary: autosomal dominant optic atrophy, autosomal recessive optic atrophy, Leber’s hereditary optic neuropathy (LHON), other mitochondrial disorders, Behr’s syndrome
- Macular dystrophies
- Deficiency: tobacco-alcohol amblyopia; vitamin B<sub>12</sub> deficiency
- Drug-induced: e.g. ethambutol, isoniazid, chloroquine
- Glaucoma

**Cross References**

Disc swelling; Foster Kennedy syndrome; Papilloedema; Temporal pallor
Optokinetic Nystagmus (OKN), Optokinetic Response

Optokinetic nystagmus (OKN) is familiar to anyone who has watched a railway passenger observing passing telegraph poles from the window of a moving train: OKN is an involuntary rhythmic eye movement induced by observing moving stimuli. In clinical practice a striped drum serves to test both visual pursuit and saccades. Rotation of the stripe to the left produces leftward pursuit, followed by a compensatory saccade to the right, followed by pursuit to the left of the next stripe, with another compensatory saccade, and so on. Hence, OKN is a physiological nystagmus.

Parietal hemisphere lesions (vascular or neoplastic) typically impair OKN. Testing for OKN may be useful in patients with suspected hysterical visual loss, since OKN cannot occur unless visual function is present; the response is lost in blindness. An internuclear ophthalmoplegia may be made more evident by testing OKN.

Cross References
Cortical blindness; Internuclear ophthalmoplegia (INO); Nystagmus; Saccades; Vestibulo-ocular reflexes

Orator’s Hand
- see BENEDICTION HAND

Oro-facial Dyspraxia
Orofacial dyspraxia, or buccofacial dyspraxia, is an inability to make voluntary, learned, movements with the oro-facial musculature, such as blowing out a match, kissing, and licking the lips. Recognized causes of oro-facial dyspraxia include

- a transient accompaniment of Broca’s aphasia, conduction aphasia, and transcortical motor aphasia of cerebrovascular origin;
- trauma to pre-Rolandic area just above the Sylvian fissure;
- in some patients with primary non-fluent aphasia; a related but distinct condition of ‘progressive loss of speech output with orofacial dyspraxia’ has also been described.

Clinical and imaging studies show a strong correlation between oro-facial dyspraxia and lesions in the frontal operculum; it may also occur with subcortical lesions involving periventricular and/or peristriatal white matter as well as the basal ganglia.

Reference

Cross Reference
Apraxia

Oromandibular Dystonia
Oromandibular dystonia, including platysma, may occur spontaneously or emerge with levodopa treatment in some patients with multiple system atrophy (MSA-P type usually), resembling risus sardonicus.
Orthostatic Hypotension

Orthostatic hypotension or postural hypotension is the finding of a persistent drop in blood pressure (BP) on standing, defined as a fall in systolic BP below 20 mmHg and diastolic BP below 10 mmHg of baseline within 3 min of adopting the upright position. Normally there is a drop in blood pressure of lesser magnitude on standing but this is usually quickly compensated for by the baroreceptor reflex. Measuring blood pressure automatically by passive head-up tilt testing (tilt table) is also helpful in diagnosing orthostatic hypotension if the active standing test is negative, and the history is suggestive, or in patients with motor impairment.

Symptoms which may be associated with orthostatic hypotension include exercise-induced or postprandial light-headedness, transient visual loss (usually bilateral), blackouts (syncope), and pain in a ‘coathanger’ distribution across the shoulders. There may be supine hypertension and reversal of the normal circadian blood pressure rhythm (normally lower at night), with an increased frequency of micturition at night. Other features of autonomic dysfunction may be present, including dry eyes and dry mouth (xerophthalmia, xerostomia), a tendency to constipation, and lack of penile erections.

Orthostatic hypotension may be found in:

- Pure autonomic failure (PAF)
- Neurodegenerative disorders such as multiple system atrophy, Parkinson’s disease, dementia with Lewy bodies
- Phaeochromocytoma
- Other causes of autonomic neuropathy (e.g. Guillain–Barré syndrome, amyloidosis)

However, the most common cause of orthostatic hypotension in hospital practice is probably dehydration or overzealous treatment with antihypertensive or diuretic agents.

Management of orthostatic hypotension consists of education on factors that influence blood pressure. Non-pharmacological approaches include increased salt and water intake, head-up bed tilt, and wearing elastic stockings or a G-suit. Pharmacological therapies include fludrocortisone (first line), and midodrine, ephedrine, or dihydroxyphenylserine (second line). Supine hypertension may also require treatment.

Reference

Cross References
Neuropathy; Parkinsonism; Xerophthalmia, Xerostomia
Oscillopsia

Oscillopsia is an illusory movement of the environment due to excessive slip of images on the retina ('retinal slip') during active or passive head movement, producing a complaint of blurring, jumping, or oscillation of the visual representation of the environment. Oscillopsia is most often due to acquired bilateral loss of vestibular function (loss of the vestibulo-ocular reflexes). This may be tested for clinically by: recording visual acuity whilst the head is passively shaken horizontally, a drop of three to seven lines of acuity versus performance with the head still suggesting loss of VOR (the dynamic illegible E test); and by observing the optic disc with an ophthalmoscope as the head is gently shaken, the disc moving with the head if VOR are lost. Vestibular testing will also demonstrate bilateral loss of vestibular function.

Recognized causes of oscillopsia include

- acquired nystagmus, e.g. pendular nystagmus;
- superior oblique myokymia;
- other ocular oscillations.

Oscillopsia does not occur in congenital nystagmus, nor in opsoclonus, presumably due to the operation of the visual suppression mechanism which normally operates during saccadic eye movements.

Oscillopsia may be treated with clonazepam.

References

Oscillucusis

Oscillucusis is an abnormal perception of an oscillation in the intensity of ambient sounds, which may occur during a migraine attack.

Reference

Osmophobia

Osmophobia, an aversion to smells, may form part of a migraine attack, along with photophobia and phonophobia.

Cross References
Phonophobia; Photophobia

Overflow

- see DYSTONIA; SYNKINESIA, SYNKINESIS
Pagophagia
- see PICA

Palatal Reflex
- see GAG REFLEX

Palatal Tremor
Palatal tremor, also known as palatal myoclonus, is characterized by rhythmic, unilateral or bilateral, palatal contractions which continue during sleep; this may be classified as a focal myoclonic syndrome. A distinction may be made between essential and symptomatic palatal tremor, also known as primary and secondary isolated palatal tremor.

Palatal tremor may be asymptomatic or there may be a clicking sound in the inner ear, especially in essential palatal tremor. There may be associated contractions of external ocular muscles (oculopalatal myoclonus), larynx, neck, diaphragm (respiratory myoclonus, diaphragmatic flutter, or Leeuwenhoek's disease), trunk, and limbs, which may bring the palatal tremor to attention. Palatal tremor may be accompanied by pendular nystagmus and oscillopsia.

Palatal myoclonus is associated with lesions interrupting pathways between the red nucleus, inferior olivary nucleus, and dentate nucleus (Guillain–Mollaret triangle). Hypertrophy of the inferior olivary nucleus may be evident neuro-radiologically (structural or functional imaging) and pathologically. This is a consequence of a lesion in the dentato-olivary pathway which leads to transsynaptic degeneration and hypermetabolism of the olivary nucleus. Although many cases are essential/idiopathic, recognized symptomatic causes of palatal tremor include vascular lesions, trauma, neoplasia, demyelination, epilepsy, and, rarely, adult-onset Alexander's disease.

Drug treatment of palatal tremor is often unsuccessful, although reports of benefit with 5-hydroxytryptophan, carbamazepine, sodium valproate, clonazepam, baclofen, and even sumatriptan have appeared. Botulinum toxin injections may also help.

Reference

Cross References
Eight-and-a-half syndrome; Myoclonus; Nystagmus; Oscillopsia; Tinnitus; Tremor

Palilalia
Palilalia is a disorder of articulation characterized by the involuntary repetition of syllables within a word, whole words, or phrases, hence a reiterative speech
disorder. The term stutter may be used for repetition of single syllables, and the term palilalia has sometimes been used for the repetition of phrases, to distinguish from palilalia. These phenomena may be encountered in:

- Parkinson’s disease (along with bradylalia, slowness of speech)
- Progressive supranuclear palsy
- Tourette syndrome (along with vocal and motor tics)
- Pick’s disease, as part of the so-called PES syndrome (palilalia, echolalia, stereotypy) or the PEMA syndrome (palilalia, echolalia, mutism, amimia)
- Late stages of Alzheimer’s disease
- Postencephalitic parkinsonism (von Economo’s disease)
- Fahr’s disease (bilateral basal ganglia calcification)
- Thalamic/midbrain infarcts
- Normal finding in children below the age of about 6 years

In pathological states, palilalia may reflect difficulty in set shifting, as seen in frontal lobe (frontal convexity) syndromes.

References


Cross References
Bradylalia; Echolalia; Frontal lobe syndromes; Hypomimia; Mutism; Parkinsonism; Stereotypy; Stutter; Tic

**Palilgia**
- see PALILALIA

**Palinacusis**
Palinacusis, or palinacousis, is the persistence of prior auditory perception. Although sometimes classified as an illusory experience, musical hallucinations may occur concurrently. The symptom may be related to seizures of temporal lobe origin.

Reference

Cross References
Hallucination; Illusion

**Palinopsia**
Palinopsia is an illusory visual phenomenon characterized by the persistence or recurrence of visual images immediately after the stimulus has been removed, hence visual perseveration. This is distinct from the physiological after-image. It may be associated with polyopia. The description of the symptom may lead to it being mistaken for diplopia (‘pseudodiplopia’).

Palinopsia occurs most frequently in the context of a left homonymous hemianopia, secondary to right occipitotemporal or occipitoparietal lesions: these may be vascular, neoplastic, metabolic, ictal, or drug- or toxin-induced (e.g.
carbon monoxide poisoning). It has also been described with retinal and optic nerve disease and occasionally in normal individuals.

References
Metz RJ, Pieri V, Diederich NJ. Object-specific and “side inversed” palinopsia limited to the hemianopic field in occipital infarction. *Journal of Neurology* 2006; 253(suppl 2): II/64 (abstract P253).

Cross References
Hemianopia; Illusion; Perseveration; Polyopia; Visual perseveration

Pallaesthesia
Pallaesthesia is the appreciation of vibration sensation; its loss may be described as pallanaethesia.

Cross Reference
Vibration

Palmaris Brevis Sign
Palmaris brevis sign may be useful in localizing the site of an ulnar nerve lesion. Innervated by the superficial ‘sensory’ division of the ulnar nerve in the wrist (distal canal of Guyon), contraction of the palmaris brevis muscle may be evident with compressive lesions of the deep motor branch of the ulnar nerve which cause intrinsic hand muscle weakness but no sensory loss (‘Ramsay Hunt syndrome’): ask the patient to ‘contract’ the hypothenar eminence with the fifth digit forcibly abducted and look for skin corrugation. In sensory superficial division ulnar nerve lesions, this sign is lost.

References

Palmomental Reflex
The palmomental reflex consists of contraction of the mentalis muscle induced by stroking the ipsilateral palm with a blunt object. It may indicate damage to the contralateral paracentral cortex or its connections, but since it is observed in about one quarter of normal adults and is very common in the normal elderly, and may occur in other conditions, both its sensitivity and specificity are low. It may be considered a frontal release sign or primitive reflex, but is less specific than the grasp reflex. Induction of the reflex by stimulation of areas other than the palm is more likely to be associated with cerebral damage.
Pandysautonomia

References

Cross References
Age-related signs; Frontal release signs

Pandysautonomia
Pandysautonomia is characterized by pre and postganglionic lesions of both the sympathetic and parasympathetic pathways. This may be

- Congenital
- Acquired:
  - acute (e.g. after a viral infection such as infectious mononucleosis)
  - subacute (e.g. the ‘autonomic-only’ form of Guillain–Barré syndrome)
  - chronic (e.g. pure autonomic failure, multiple system atrophy, certain hereditary neuropathies)

Clinical features include

- Visual blurring; pupillary areflexia
- Orthostatic hypotension
- Cardiac arrhythmia
- Abdominal pain, diarrhoea, vomiting, constipation, ileus, pseudo-obstruction

Response to intravenous immunoglobulin has been reported in idiopathic pandysautonomia.

Reference

Papilloedema
Papilloedema is swelling (oedema) of the optic nerve head due to raised intracranial pressure (cf. other causes of disc swelling, which may cause pseudopapilloedema). A number of stages of papilloedema are described: in the acute stage, the only findings may be oedema at the superior and inferior poles of the disc, absence of spontaneous venous pulsation, and enlargement of the blind spot. As papilloedema progresses the whole disc is involved and splinter haemorrhages may be evident at the disc margin. These early stages may be asymptomatic or may be associated with transient losses of vision (obscurations), often provoked by activities or movements which further raise intracranial pressure, thus compromising retinal perfusion pressure. Enlargement of the blind spot and constriction of the visual field may be evident, but visual acuity is often unimpaired (cf. disc swelling due to papillitis). Chronic papilloedema produces gliosis of the optic nerve head and eventually optic atrophy (‘sequential optic atrophy’) with nerve fibre damage and permanent visual field defects.

Cross References
Blind spot; Disc swelling; Obscurations; Optic atrophy; Pseudopapilloedema; Retinal venous pulsation; Scotoma
Paraesthesia

Paraesthesia is an abnormal sensation, often described as a tingling sensation, or likened to ‘pins and needles’ or electricity, pricking, or even crawling (formication), i.e. positive sensory symptoms. The sensation is not pleasant but nor is it painful (cf. dysesthesia). Some patients may describe this sensation as ‘numbness’ or ‘deadness’, in which case care needs to be taken to differentiate it from anaesthesia (i.e. a negative phenomenon). Some authorities reserve the term for spontaneous rather than evoked positive sensory phenomena, as a distinction from dysesthesia.

Paraesthesia is a feature of neuropathy and may occur in the distribution of a compressed or entrapped nerve, perhaps reflecting the mechanosensitivity of nerves in this situation (e.g. Phalen’s sign, Tinel’s sign). Paraesthesia is a more reliable indicator of the diagnosis of neuropathy than pain. Paraesthesia may also be provoked by hyperventilation (especially perioral, hands, and feet [acroparaesthesia]). Central lesions may also produce paraesthesia (e.g. Lhermitte’s sign).

Reference

Cross References
Anaesthesia; Dysesthesia; Lhermitte’s sign; Phalen’s sign; Tinel’s sign
Paragrammatism
Paragrammatism is the substitution of morphological elements and function words in the context of fluent speech (e.g. Wernicke’s aphasia), as differentiated from agrammatism, the omission of function words and bound morphemes in non-fluent speech (e.g. Broca’s aphasia).

Cross References
Agrammatism; Aphasia; Broca’s aphasia; Wernicke’s aphasia

Paragraphia
- see AGRAPHIA

Parakinesia, Parakinesis
These terms have been used in different ways by different authors, to describe

- a volitional purposeful act designed to camouflage, mask, or draw attention away from an involuntary movement, such as chorea;
- strange movements of presumed psychogenic origin. It should be remembered that many movements previously thought to conform to this definition have subsequently been recognized to have an organic basis (e.g. klazomania).

The terms are now seldom used.

Cross References
Chorea, Choreoathetosis; Dyskinesia; Klazomania

Paralexia
- see ALEXIA

Paralogia
- see GANSER PHENOMENON

Paralysis
Paralysis is a total loss of power to move a body part; equivalent to the suffix -plegia. The use of the word has not been entirely consistent, for example, paralysis agitans originally used by James Parkinson to describe the disease which now bears his name. The periodic paralyses are a group of conditions characterized by episodic muscular weakness and stiffness (myotonia) associated with mutations in the skeletal muscle voltage-gated sodium and calcium ion channel genes (channelopathies).

Cross References
Myotonia; Plegia

Paramnesia
Paramnesia is recalling as memories things which have not in fact taken place, hence a distortion of episodic or autobiographical memory. This may be neurological or psychiatric in origin. Relation of paramnesias as the truth occurs in confabulation.

Cross References
Amnesia; Confabulation; Reduplicative paramnesia
Paramyotonia

Paramyotonia is similar to myotonia in that muscle does not relax normally following contraction (voluntary, percussion), which may prompt a complaint of muscle aching or stiffness, but differs in that repetitive muscle use (e.g. exercise) accentuates the problem, leading to an increased delay in muscle relaxation (worsening stiffness). For example, repeated forced voluntary eyelid closure in a patient with paramyotonia may, after several attempts, lead to a failure of voluntary eyelid opening, the eyes remaining closed for a minute or so. Paramyotonia particularly affects the face and forearms. This type of muscle stiffness may also be sensitive to temperature, being made worse by cooling which may also provoke muscle weakness. Weakness may outlast exposure to cold by several hours. Neurophysiological studies may assist in the diagnosis of paramyotonia. During the delayed muscle relaxation, electrical activity is not prominent, and after muscle cooling the resting muscle membrane potential may be reduced from around the normal $-80$ to $-40$ mV, at which point muscle fibres are inexcitable (contracture).

Paramyotonia congenita (Eulenburg’s disease) is a channelopathy with mutations affecting the $\alpha$-subunit of the sodium channel (SCN4A). Mutations in the same gene have been documented in hyperkalaemic periodic paralysis and K$^+$-aggravated myotonia.

Symptomatic treatment with membrane-stabilizing agents like mexiletine and tocainide or with the carbonic anhydrase inhibitor acetazolamide might be tried. Precautions are necessary during general anaesthesia because of the risk of diaphragm myotonia.

References

Cross References
Contracture; Myotonia; Paralysis; Warm-up phenomenon

Paraparesis

Paraparesis is a weakness of the lower limbs, short of complete weakness (paraplegia). This may result from lesions anywhere from cerebral cortex (frontal, parasagittal lesions) to peripheral nerves, producing either an upper motor neurone (spastic) or lower motor neurone (flaccid) picture. A spinal cord lesion (myelopathy) is probably the most common cause. Paraparesis may be symmetrical or asymmetrical. Recognized causes of paraparesis include

- **Upper motor neurone lesions:**
  - Traumatic section of the cord;
  - Cord compression from intrinsic or extrinsic mass lesion, e.g. tumour, metastasis, abscess, empyema, haematoma (epidural, subdural);
  - Inflammatory lesions: acute transverse myelitis of viral origin, multiple sclerosis, neuromyelitis optica (Devic’s syndrome), systemic lupus erythematosus, Behçet’s disease, giant cell arteritis (rare);
Paraphasia

Structural lesions: tethered cord syndrome, arteriovenous malformation;
Metabolic: hereditary spastic paraplegia (HSP), adrenoleucodystrophy (X-ALD), subacute combined degeneration of the cord (usually mild).

- **Lower motor neurone lesions:**
  Acute or chronic neuropathies (Guillain–Barré syndrome, chronic inflammatory demyelinating polyradiculoneuropathy).

**Cross References**
Flaccidity; Myelopathy; Paraplegia; Spasticity

**Paraphasia**
Paraphasias are a feature of aphasias (disorders of language), particularly (but not exclusively) fluent aphasias resulting from posterior dominant temporal lobe lesions (cf. anterior lesions which tend to produce non-fluent aphasias with agrammatism). Paraphasias refer to a range of speech output errors, both phonological and lexical, including substitution, addition, duplication, omission, and transposition of linguistic units, affecting letters within words, letters within syllables, or words within sentences. Paraphasic errors may be categorized as:

- **Phonemic or literal:**
  Errors involve individual phonemes; impaired phonology (i.e. sound based) causing approximations to real words; non-words resulting from phonemic paraphasia may be referred to as neologisms. Phonemic paraphasias may be encountered in Broca’s aphasia and conduction aphasia, when the patient may recognize them to be errors, and Wernicke’s aphasia.

- **Formal:**
  Target word is replaced by another word that is phonemically similar.

- **Morphemic:**
  Errors involving word stems, suffixes, prefixes, inflections, and other parts of words.

- **Verbal:**
  Errors involving whole words. These may be further classified as:
  - **Semantic or categoric:** substitution of a different exemplar from the same category (e.g. ‘orange’ for ‘apple’; paradigmatic) or of a thematically related word (e.g. ‘sit’ for ‘chair’; syntagmatic). Verbal paraphasias showing both semantic and phonemic resemblance to the target word are called *mixed* errors. These types may be observed in patients with Wernicke’s aphasia, who often seem unaware of their paraphasias due to a failure of self-monitoring of output.

**Reference**
Paraplegia

Paraplegia is a total weakness (paralysis) of the lower limbs (cf. paraparesis). This may result from lower motor neurone lesions involving multiple nerve roots and/or peripheral nerves (e.g. paraparetic Guillain–Barré syndrome) producing a flaccid, areflexic paraplegia; but more commonly it is due to upper motor neurone lesions interrupting corticospinal pathways (corticospinal tract, vestibulospinal tract, reticulospinal tracts, and other extrapyramidal pathways), most usually in the spinal cord. The latter may acutely produce a flaccid areflexic picture (‘spinal shock’), but later this develops into an upper motor neurone syndrome (hypertonia, clonus, hyperreflexia, loss of superficial reflexes [e.g. abdominal, cremasteric reflexes] and Babinski’s sign) with possible lower motor neurone signs at the level of the lesion; bladder involvement is common (urinary retention). Because of the difficulty in distinguishing whether an acute paraplegia is of LMN or UMN origin, imaging to exclude potentially reversible cord compression is mandatory. Recognized causes of paraplegia of upper motor neurone origin include

- traumatic section of the cord;
- cord compression;
- inflammatory lesions: acute transverse myelitis of viral origin, multiple sclerosis, neuromyelitis optica (Devic’s syndrome);
- ischaemic lesions; anterior spinal artery syndrome, venous infarction of the cord.

In paraplegia of upper motor neurone origin, enhanced flexion defence reflexes (‘flexor spasms’) may occur, producing hip and knee flexion, ankle and toe dorsiflexion. Eventually such flexor responses may become a fixed flexion deformity with secondary contractures (‘paraplegia in flexion’). Prevention of this situation may be possible by avoiding spasms, which are often provoked by skin irritation or ulceration, bowel constipation, bladder infection, and poor nutrition. Physiotherapy and pharmacotherapy with agents such as baclofen, dantrolene, and tizanidine may be used; botulinum toxin injections may be helpful for focal spasticity. ‘Paraplegia in extension’, with extension at the hip and knee, may be seen with incomplete or high spinal cord lesions.

Reference

Cross References
Abdominal reflexes; Areflexia; Babinski’s sign (1); Clonus; Contracture; Cremasteric reflex; Flaccidity; Hyperreflexia; Hypertonia, Hypertonus; Lower motor neurone (LMN) syndrome; Myelopathy; Paraparesis; Spasticity; Upper motor neurone (UMN) syndrome; Urinary retention

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**Parapraxia, Parapraxis**

Although this term may be used in common parlance as synonymous with a ‘Freudian slip’, in neurological practice it has a different meaning, referring to one of the cardinal symptoms of ideomotor apraxia: a combination of deficient action selection with errors of sequencing of actions and spatial orientation errors. Parapraxic errors include

- **Perseveration**: repetition of movements;
- **Substitution**: of one movement for another;
- **Surplus movements**;
- **Verbal overflow**: explaining a movement rather than performing it;
- **Omission**: incomplete movements;
- **Conduit d’approche**: several attempts to perform the correct movement;
- **Body part as object**.

**Cross References**

Apraxia; Body part as object; *Conduit d’approche*; Perseveration

**Paratonia**

- see **GEGENHALTEN**

**Paresis**

Paresis denotes a weakness which is less than total paralysis (-plegia), which may be of upper or lower motor neurone origin. Various prefixes denote the location of such weakness, e.g. hemiparesis, monoparesis, ophthalmoparesis, paraparesis, quadriparesis.

Since localized pain may inhibit voluntary muscular exertion, apparent weakness in such circumstances may be labelled ‘algesic pseudoparesis’.

**Cross References**

Lower motor neurone (LMN) syndrome; Paralysis; Plegia; Upper motor neurone (UMN) syndrome; Weakness

**Parinaud’s Syndrome**

Parinaud’s syndrome, also sometimes known as the dorsal midbrain syndrome, the periaqueductal grey matter syndrome, or the pretectal syndrome, consists of:

- **Eye movements**:
  
  - Paralysis of vertical gaze, especially upgaze: Bell’s phenomenon may be spared
  - Loss of convergence; convergence spasm may cause slow abduction (‘midbrain pseudo-sixth’)
  - Skew deviation
  - Convergence-retraction nystagmus (Körber–Salus–Elschnig syndrome); sometimes downbeat nystagmus

- **Eyelids**:
  
  - Lid retraction (Collier’s ‘tucked lid’ sign) or ptosis (ventral extension of lesion)
Pupils:

Mydriasis

This constellation of signs results from dorsal midbrain lesions, such as pineal tumours, which affect the pretectum and posterior commissure and so interfere with conjugate eye movements in the vertical plane. The key anatomical substrates, damage to which causes the syndrome, are probably the interstitial nucleus of Cajal and the nucleus of the posterior commissure and their projections.

References

Cross References
Collier’s sign; Light-near pupillary dissociation; Nystagmus; Supranuclear gaze palsy

Parkinsonism
Parkinsonism is a clinical syndrome characterized by the presence of some or all of the following features; there is overlap with so-called akinetic-rigid syndromes in which these features predominate:

- Akinesia, hypokinesia (sine qua non)
- Rigidity: consistent (leadpipe) or jerky (cogwheeling; Negro’s sign)
- Bradykinesia
- Tremor, usually at rest, of frequency 3.5–7.0 Hz, ‘pill rolling’ type; there may sometimes be an additional action component to the tremor, and very occasionally there is exclusively an action tremor. ‘Re-emergent tremor’ is also described
- Stooped posture: forward flexion of trunk, flexion of knees, elbows; ‘simian posture’
- Impaired postural reflexes, with or without a history of falls; propulsion, retropulsion
- Mask-like facies, poverty of spontaneous facial expression (hypomimia)
- Reduced blink rate (this may be a particular feature of progressive supranuclear palsy)
- Hypophonic, monotonic voice (hypokinetic dysarthria)
- Widened palpebral fissure (Stellwag’s sign)
- Hypometria
- Seborrhoea
- Sialorrhoea
- Festinant (shuffling) gait
- Micrographia
- Dystonic postures, e.g. striatal toe
- Apraxia
• Akathisia
• Cognitive impairment (usually of frontal–subcortical type)
• Hallucinations: minor (anwesenheit; passage type), or formed, visual > auditory. Insight into the non-reality of these experiences may be retained, hence ‘pseudohallucinations’
• Autonomic dysfunction, especially orthostatic hypotension

Conventionally parkinsonism is viewed as a disorder of the extrapyramidal system producing ‘extrapyramidal signs’, although this term has limitations: despite the fact that some of the cardinal features of parkinsonism (bradykinesia, rigidity, postural instability, tremor) result from pathology in the basal ganglia, particularly affecting dopaminergic pathways, other features may reflect cortical involvement, at least in part (e.g. apraxia, micrographia).

The incidence of parkinsonism increases dramatically with age; it is also associated with an increased risk of death, particularly in the presence of a gait disturbance.

The differential diagnosis of parkinsonism is broad, and includes

• Idiopathic Parkinson’s disease
• Multiple system atrophy
• Progressive supranuclear palsy (Steele–Richardson–Olszewski syndrome)
• Corticobasal degeneration, cortical basal ganglionic degeneration
• Drug-induced parkinsonism (e.g. neuroleptics, MPTP)
• Toxin-induced parkinsonism (e.g. carbon monoxide, manganese)
• Wilson’s disease (hepatolenticular degeneration); non-Wilsonian hepatocerebral degeneration
• Dementia with Lewy bodies
• Neuroleptic malignant syndrome
• Normal pressure hydrocephalus
• ‘Arteriosclerotic parkinsonism’, resulting from multiple subcortical infarcts
• Huntington’s disease, especially juvenile onset (Westphal variant)
• Postencephalitic parkinsonism (encephalitis lethargica, von Economo’s disease)
• Dementia pugilistica, posttraumatic parkinsonism
• Systemic lupus erythematosus
• Sjögren’s syndrome
• Hypoparathyroidism
• Parkinsonism–dementia complex of Guam

Obsessive slowness also enters the differential diagnosis but typical parkinsonian features (akinesia, rigidity) are not present in this condition.

It is crucial not to miss the diagnosis of Wilson’s disease, although rare, since in the early stages this disorder is reversible with copper chelation therapy; hence copper and caeruloplasmin should be checked in all patients with young-onset (under age 50) parkinsonism (and dystonia).

Response to levodopa therapy is only reliably seen in idiopathic Parkinson’s disease, although some patients with multiple system atrophy or progressive supranuclear palsy may benefit. The features particularly responsive in Parkinson’s disease are bradykinesia and rigidity; tremor is less reliably helped.
Parry–Romberg Syndrome

References

Cross References
Apraxia; Blinking; Bradykinesia; Dysarthria; Dystonia; Hypokinesia; Hypomimia; Hypophonia; Mask-like facies; Micrographia; Orthostatic hypotension; Postural reflexes; Rigidity; Seborrhoea; Sialorrhoea; Striatal toe; Supranuclear gaze palsy; Tremor

Parosmia
Parosmia is a false smell, i.e. the subjective sensation of a smell which does not exist (i.e. an hallucination). Such smells are usually unpleasant (cacosmia), may be associated with a disagreeable taste (cacoguesia), and may be difficult for the patient to define. Causes include purulent nasal infections or sinusitis and partial recovery following transection of olfactory nerve fibres after head injury. Transient parosmia may presage epileptic seizures of temporal lobe cortical origin (olfactory aura), particularly involving the medial (uncal) region. The symptom may also be common amongst the normal population.

Reference

Cross References
Aura; Seizures

Parry–Romberg Syndrome
Hemifacial atrophy is thinning of subcutaneous tissues on one side of the face; it may also involve muscle and bone (causing enophthalmos), and sometimes brain, in which case neurological features (hemiparesis, hemianopia, focal epileptic seizures, cognitive impairment) may also be present.

The clinical heterogeneity of hemifacial atrophy probably reflects patho-
genetic heterogeneity. The syndrome may result from maldevelopment of autonomic innervation or vascular supply, or as an acquired feature following trauma, or a consequence of linear scleroderma (morphoea), in which case a coup de sabre may be seen.

References

Cross References
Coup de sabre; Enophthalmos; Hemianopia; Hemiparesis
Past-Pointing
- see DYSMETRIA

Patellar Reflex
- see REFLEXES

Pathological Crying, Pathological Laughter
Pathological laughter and pathological crying (PLC), or forced laughter and crying, also referred to as involuntary emotional expression disorder, have been defined as reflecting an incongruence of mood (subjective feeling) and expression or affect (‘objective’, observed), such that patients laugh involuntarily though not happy, or cry though not sad. There may be a sense that the patient is struggling against these displays of emotion, in contrast to the situation in other forms of emotional lability where there is said to be congruence of mood and affect, although sudden fluctuations and exaggerated emotional expression are common to both, suggesting a degree of overlap.

PLC are ascribed to a loss (release) of the normal inhibition of the motor component of facial expression (i.e. cortical–subcortical disinhibition). PLC may occur in the context of a pseudobulbar palsy (‘pseudobulbar affect’) but not invariably so. PLC have been reported in:

- Multiple sclerosis: crying > laughing; related to intellectual impairment (more extensive brain involvement, but not brainstem);
- Alzheimer’s disease;
- Stroke: PLC may be the harbinger of brainstem stroke or a feature of anterior choroidal artery territory infarctions; rarely a feature of TIAs;
- Motor neurone disease;
- Head injury;
- Gelastic epilepsy.

A Pathological Laughter and Crying Scale has been developed. Suggested treatments for PLC include amitriptyline, levodopa, amantadine, and serotonin-reuptake inhibitors such as fluoxetine and citalopram.

References

Cross References
Automatism; Emotionalism, Emotional lability; Pseudobulbar palsy

Peduncular Hallucinosis
Peduncular hallucinosis is a rare syndrome characterized by hallucinations and brainstem symptoms. Hallucinations are vivid and naturalistic. Brainstem findings include oculomotor disturbances, dysarthria, ataxia, and impaired arousal.
Episodic memory impairments also occur. Pathology may be in midbrain, thalamus, and pons.

**Reference**

**Cross Reference**
Hallucination

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**Peek Sign**
One of the eye signs of myasthenia gravis: on attempted forced eye closure, orbicularis oculi may fatigue such that the patient ‘peeks’ through the partially open palpebral fissure.

**Pelopsia, Peliopsia**
Peliopsia or pelopsia is a form of metamorphopsia characterized by the misperception of objects as closer to the observer than they really are (cf. porropsia, telopsia).

**Cross References**
Metamorphopsia; Porropsia

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**Pelvic Thrusting**
Pelvic thrusting may be a feature of epileptic seizures of frontal lobe origin; occasionally it may occur in temporal lobe seizures. Pelvic thrusting also occurs in pseudoseizures, particularly those of the ‘thrashing’ variety. Choreiform disorders may involve the pelvic region causing thrusting or rocking movements.

**Reference**

**Cross References**
Automatism; Chorea, Choreoathetosis; Seizure

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**Pendular Nystagmus**
Pendular or undulatory nystagmus is characterized by eye movements which are more or less equal in amplitude and velocity (sinusoidal oscillations) about a central (null) point. In acquired causes such as multiple sclerosis, this may produce oscillopsia and blurred vision. Treatment options include gabapentin and memantine.

**Reference**

**Cross References**
Nystagmus; Oscillopsia

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**Percussion Myotonia**
Percussion myotonia is the myotonic response of a muscle to a mechanical stimulus, e.g. when struck with a tendon hammer. For example, a blow to the thenar eminence may produce involuntary and sustained flexion of the thumb. This
response, which may be seen in myotonic dystrophy, reflects the impaired muscle relaxation which characterizes myotonia.

**Cross Reference**
Myotonia

**Periodic Alternating Nystagmus**
Periodic alternating nystagmus is a horizontal jerk nystagmus, which damps or stops for a few seconds and then reverses direction. Eye movements may need to be observed for up to 5 min to see the whole cycle. Periodic alternating nystagmus may be congenital or acquired, if the latter then its localizing value is similar to that of downbeat nystagmus (with which it may coexist), especially for lesions at the cervico-medullary junction (e.g. Chiari malformation). Treatment of the associated lesion may be undertaken, otherwise periodic alternating nystagmus usually responds to baclofen, hence the importance of correctly identifying this particular form of nystagmus.

**Reference**

**Cross Reference**
Nystagmus

**Periodic Respiration**
Periodic respiration is a cyclical waxing and waning of the depth and rate of breathing (Cheyne–Stokes breathing or respiration), over about 2 min, the crescendo–decrescendo sequence being separated by central apnoeas. A so-called variant Cheyne–Stokes pattern has hypopnoeas rather than apnoeas.

Periodic respiration may be observed in unconscious patients with lesions of the deep cerebral hemispheres, diencephalon, or upper pons, or with central or tonsillar brain herniation; it has also been reported in multiple system atrophy. Prolonged circulatory time (congestive heart failure) and hypoxaemia (e.g. at altitude) may also cause periodic respiration, but with a shorter cycle.

**Reference**

**Cross References**
Coma

**Perseveration**
Perseveration refers to any continuation or recurrence of activity without appropriate stimulus (cf. intrusions). Perseverations may be repeated motor behaviours (e.g. drawing, writing) or speech. These are viewed as a failure to inhibit a previous response pattern. Sensory perseveration is also described, e.g. palinopsia in the visual system. A number of varieties of perseveration have been described, associated with lesions in different areas of the brain:

- ‘*Stuck-in-set*’:
  
  Inappropriate maintenance of a current category or framework; thought to reflect a deficit in executive function; associated with frontal lobe (especially frontal convexity) damage, which is associated with
an inert, apathetic pattern of behaviour, rather than the disinhibited pattern associated with orbitofrontal damage.

- ‘Recurrence’:
  Unintentional repetition of a previous response to a subsequent stimulus; thought to represent an abnormal post-facilitation of a memory trace; associated with posterior left (dominant) hemisphere damage; commonly seen in aphasics, Alzheimer’s disease; this overlaps with ‘intrusions’.

- ‘Continuous’:
  Inappropriate prolongation or repetition of a current behaviour without interruption; thought to represent a deficit of motor output; associated with basal ganglia damage.

References

Cross References
Aphasia; Dysexecutive syndrome; Frontal lobe syndromes; Intrusion; Logoclonia; Palinopsia

**Personification of Paralyzed Limbs**
Critchley drew attention to the tendency observed in some hemiplegic patients to give their paralyzed limbs a name or nickname and to invest them with a personality or identity of their own. This sometimes follows a period of anosognosia and may coexist with a degree of anosodiaphoria; it is much more commonly seen with left hemiplegia. A similar phenomenon may occur with amputated limbs, and it has been reported in a functional limb weakness.

References

Cross References
Anosodiaphoria; Anosognosia

**Pes Cavus**
Pes cavus is a high-arched foot due to equinus (plantar flexion) deformity of the first ray, with secondary changes in the other rays (i.e. deformity is more evident on the medial side of the foot in most cases). This may be due to imbalance of muscular forces during development (e.g. strong peroneus longus, weak peroneus brevis and tibialis anterior, although the precise pattern may differ with cause), which may be a consequence of neurological disease. Hammer toes may also be present. Pes cavus may be associated with disease of genetic origin, e.g. hereditary motor and sensory neuropathy (HMSN, Charcot–Marie–Tooth syndrome), hereditary spastic paraparesis, Friedreich’s ataxia, Marfan’s syndrome;
or be due to an early neurological insult, e.g. cerebral palsy, paralytic poliomyelitis. Familial pes cavus without other neurological signs has also been reported (a *forme fruste* of HMSN?). Surgical treatment of pes cavus may be necessary, especially if there are secondary deformities causing pain, skin breakdown, or gait problems.

**Cross References**
Claw foot; Hammer toes

**Petite Madeleines Phenomenon**
- see PROUST PHENOMENON

**Phalen’s Sign**
Phalen’s sign is present when tingling (paraesthesia) is experienced in the distribution of the median nerve when the wrist is held in forced flexion (90° for 30–60 s; Phalen’s manoeuvre). Patients may volunteer that they experience such symptoms when carrying heavy items such as shopping bags which puts the hand in a similar posture. Hyperextension of the wrist (‘reverse Phalen’s manoeuvre’) may also reproduce symptoms. These are signs of compression of the median nerve at the wrist (carpal tunnel syndrome). Like other provocative tests (e.g. Tinel’s sign), the sensitivity and specificity of Phalen’s sign for this diagnosis are variable (10–91% and 33–86%).

The pathophysiology of Phalen’s sign is probably the lower threshold of injured nerves to mechanical stimuli, as for Tinel’s sign and Lhermitte’s sign.

**References**
D’Arcy CA, McGee S. Does this patient have carpal tunnel syndrome? *JAMA* 2000; **283**: 3110–3117.

**Cross References**
Flick sign; Lhermitte’s sign; Paraesthesia; Tinel’s sign

**Phantom Alloaesthesia**
- see ALLOAESTHESIA

**Phantom Chromatopsia**
This term has been coined to refer to the complaint of patients who are blind or nearly so that a colour, usually golden or purple, enlarges to invade the entire visual field. This is presumably cortical in origin and has been described as an hallucination. ‘Phantom vision’ may describe a similar phenomenon.

**Reference**

**Cross References**
Erythropsia; ‘Monochromatopsia’; Phantom vision

**Phantom Limb**
Phantom limbs, or ghost limbs, are the subjective report of the awareness of a non-existing or deafferented body part in a mentally otherwise competent
individual. The term was coined by Weir Mitchell in the nineteenth century, but parts other than limbs (either congenitally absent or following amputation) may be affected by phantom phenomena, such as lips, tongue, nose, eye, penis, breast and nipple, teeth, and viscera. Phantom phenomena are perceived as real by the patient, may be subject to a wide range of sensations (pressure, temperature, tickle, pain), and are perceived as an integral part of the self. Such ‘limbless perception’ is thought to reflect the mental representation of body parts generated within the brain (body schema), such that perception is carried out without somatic peripheral input. Reorganization of cortical connections following amputation may explain phantom phenomena such as representation of a hand on the chest or face, for which there is also evidence from functional brain imaging.

References

Phantom Vision
This name has been given to visual hallucinations following eye enucleation, by analogy with somaesthetic sensation experienced in a phantom limb after amputation. Similar phenomena may occur after acute visual loss and may overlap with phantom chromatopsia. Unformed or simple hallucinations are more common than formed or complex hallucinations.

References

Cross References
Hallucination; Phantom chromatopsia; Phantom limb

Pharyngeal Reflex
- see GAG REFLEX

Phonagnosia
Phonagnosia is an inability to recognize familiar voices in the absence of hearing impairment, hence a form of auditory agnosia. The patient can recognize and understand words and sentences (cf. pure word deafness). Phonagnosia is the equivalent in the auditory domain of prosopagnosia in the visual domain. The neuroanatomical substrate is thought to be right parietal lobe pathology.

Cross References
Agnosia; Auditory agnosia; Prosopagnosia; Pure word deafness

Phonemic Disintegration
Phonemic disintegration refers to an impaired ability to organize phonemes, the smallest units in which spoken language may be sequentially described, resulting
in substitutions, deletions, and misorderings of phonemes. Phonemic disintegration is relatively common in aphasic disorders, including Broca’s aphasia, conduction aphasia, and transcortical motor aphasia. Isolated phonemic disintegration is rare. The neural substrate may be primary motor cortex of the left inferior precentral gyrus and subjacent white matter, with sparing of Broca’s area.

References

Cross References
Aphasia; Aphemia; Broca’s aphasia

**Phonetic Disintegration**
- see APHEMIA

**Phonophobia**
Phonophobia is a dislike, or fear, of sounds, especially loud sounds, often experienced during a migraine headache.

Cross Reference
Hyperacusis

**Phosphene**
Phosphenes are percepts in one modality induced by an inappropriate stimulus, e.g. when pressure is applied to the eyeball, the mechanical stimulus may induce the perception of light. The perception of flashes of light when the eyes are moved has been reported in optic neuritis, presumably reflecting the increased mechanosensitivity of the demyelinated optic nerve fibres; this is suggested to be the visual equivalent of Lhermitte’s sign. Eye gouging to produce phosphenes by mechanical stimulation of the retina is reported in Leber’s congenital amaurosis. Noise-induced visual phosphenes have also been reported and may be equivalent to auditory-visual synaesthesia.

References

Cross References
Auditory-visual synaesthesia; Gaze-evoked phenomena; Lhermitte’s sign; Photism; Synaesthesia

**Photism**
Photisms are transient positive visual phenomenon, such as geometrical shapes or brightly coloured spectral phenomena, occurring in the context of epilepsy, migraine, or in blind visual fields (hence overlapping with photopsia). Auditory-visual synaesthesia may also be described as sound-induced photism.
Photophobia
Photophobia is an abnormal intolerance of light, often experienced with eye pain. It is associated with a wide range of causes and may result from both peripheral and central mechanisms:

- Anterior segment eye disorders: uveitis, glaucoma, cataract;
- Vitreo-retinal disorders: retinitis pigmentosa;
- Optic neuropathies: optic neuritis;
- Intracranial disease: migraine, meningitis, and other causes of meningeal irritation, central photophobia (?thalamic lesion), dazzle;
- Physiological photophobia: sudden exposure to light after light deprivation.

Cross References
Dazzle; Meningism; Retinitis pigmentosa

Photopsia
Photopsias are simple visual hallucinations consisting of flashes of light which often occur with a visual field defect. They suggest dysfunction in the inferomedial occipital lobe, such as migraine or an epileptogenic lesion.

Cross References
Aura; Hallucination; Photism

Physical Duality
A rare somaesthetic metamorphopsia occurring as a migraine aura in which individuals feel as though they have two bodies.

Cross Reference
Metamorphopsia

Piano-Playing Fingers
- see PSEUDOATHETOSIS

Pica
Pica, or pagaphagia, is a morbid craving for unusual or unsuitable food in association with iron deficiency. It has also been reported in tuberous sclerosis. Sufferers risk infection from contaminated foods.

References

Cross Reference
Geophagia, Geophagy

Picture Sign
The 'picture sign' is present when a patient believes that individuals seen on the television screen are actually present in the home; indeed they may be reported
to emerge from the television set into the room. This may occur as part of the
cognitive disturbance of Alzheimer’s disease or dementia with Lewy bodies, or
as part of a psychotic disorder. Like the ‘mirror sign’, the ‘picture sign’ may be
classified as a misidentification phenomenon.

Cross References
‘Mirror sign’; Misidentification syndromes

‘Picture Within a Picture’ Sign
Following a right parieto-occipital infarction, a patient complained of seeing
people moving about in the left lower quadrant of the visual field whilst vision
was normal in the remainder of the visual field, a phenomenon labelled the
‘picture within a picture’ sign. This has been categorized as a visual release
hallucination.

Reference
Benegas MN, Liu GT, Volpe NJ, Galetta SL. “Picture within a picture” visual

Pied en Griffe
- see CLAW FOOT

‘Pie-in-the-Sky’ Defect
This name has sometimes been given to the superior homonymous quadrantan-
ipia ending sharply at the vertical midline due to a temporal lobe lesion
interrupting Meyer’s loop, that part of the optic radiation coursing through the
temporal lobe.

Cross Reference
Quadrantanopia

Pill Rolling
- see PARKINSONISM; TREMOR

Pinch Sign
The ‘pinch sign’, or ‘okay sign’, is an inability to make a small circle (‘form the
letter O’, divers’ okay sign) by approximating the distal phalanges of the thumb
and index finger, due to weakness of flexor digitorum profundus in the index fin-
ger and flexor pollicis longus in the thumb as a consequence of median nerve
lesions in the forearm, e.g. anterior interosseous neuropathy, pronator teres syn-
drome. This results in a pinching posture of thumb and index finger. The ‘straight
thumb sign’ may also be present.

Cross References
Froment’s sign; ‘Straight thumb sign’

Pinhole Test
Impairments in visual acuity due to refraction defects (changes in shape of the
globe or defects in the transparent media of the eye) may be improved or cor-
rected by looking through a pinhole which restricts vision to the central beam of
light.
**Plantar Response**

The plantar response is most commonly elicited by stroking the sole of the foot with a blunt object. The first response of the hallux is the critical observation, which may be facilitated by having one’s line of vision directly above the axis of the toe. The normal response after maturation of the corticospinal tracts (i.e., after about 3 years of age) is for the big toe to flex. An extensor response of the big toe in an adult (Babinski’s sign), with or without fanning (abduction) of the other toes (fan sign, *signe de l’éventail*), is a reliable sign of upper motor neurone pathology. Use of the term ‘negative Babinski’s sign’ or ‘negative Babinski response’ to mean ‘flexor plantar response’ is incorrect and should not be used. This normal plantar response is a superficial cutaneous reflex, analogous to abdominal and cremasteric reflexes, whereas the pathological response is often accompanied by activity in other flexor muscles. In some individuals the toes do not move at all, in which case the response is labelled as ‘mute’ or absent. Assessment of the response may be confounded by withdrawal of the foot in ticklish individuals. Differentiation from the striatal toe seen in parkinsonian syndromes is also important.

The plantar response may be elicited in a variety of other ways which are not in routine clinical use. Of these, perhaps the most frequently used are Chaddock’s sign (application of a stimulus in a circular direction around the external malleolus or the lateral aspect of the foot from heel to little toe) and Oppenheim’s sign (application of heavy pressure to the anterior surface of the tibia from patella to ankle). These may be helpful in ticklish patients who object to having their feet stroked. If the plantar response thus elicited is upgoing, this suggests a spread of the ‘receptive field’ of the reflex. Babinski’s sign is the earliest to occur in the presence of upper motor neurone pathology.

It is often difficult to form a definite judgment on the plantar response and reproducibility is also questionable. A study of 24 experienced clinicians invited to examine plantar responses ‘blind’ found that the interobserver percentage agreement beyond chance was on average only 16.7% (95% confidence interval [CI] 0.4–33%); intraobserver percentage agreement was a little better (average 59.6%; CI 39.6–79.6%). There remains a persistent belief, particularly amongst trainees, that an experienced neurologist can make the plantar response go whichever way s/he chooses.

**References**


**Cross References**

Abdominal reflexes; Babinski’s sign (1); Chaddock’s sign; Gordon’s sign; Oppenheim’s sign; Reflexes; Striatal toe, Upper motor neurone (UMN) syndrome

**Plegia**

Plegia means stillness, implying a complete weakness (or paralysis in common parlance), as in monoplegia, diplegia, ophthalmoplegia, paraplegia, quadriplegia, and cardioplegia. Hence plegia is a more severe weakness than paresis.
Pleurothotonus

Pleurothotonus, or Pisa syndrome, is a truncal dystonia characterized by involuntary side flexion of the head and neck, which may occur as an adverse effect of neuroleptics, antiemetics, valproate, atypical antipsychotics, and cholinesterase inhibitors.

Cross Reference
Dystonia

Plexopathy

Lesions confined to the brachial, lumbar, or sacral plexi may produce a constellation of motor and sensory signs (weakness, reflex diminution or loss, sensory loss) which cannot be ascribed to single or multiple roots (radiculopathy) or peripheral nerves (neuropathy). Lesions may involve the whole plexus (panplexopathy):

- **Brachial:** C5–T1
- **Lumbar:** L2–L4
- **Sacral:** L5–S3

or be partial, e.g. upper trunk of brachial plexus (C5–C6), producing ‘waiter’s tip’ posture (as for C5/C6 root avulsion); lower trunk of brachial plexus (C8–T1; as for C8/T1 root avulsion).

Neurophysiological studies may be helpful in distinguishing plexopathy from radiculopathy: sensory nerve action potentials (SNAPs) are reduced or absent in plexopathies because the lesion is located distal to the dorsal root ganglion (DRG), whereas SNAPs are normal in radiculopathies because the lesion is proximal to the DRG. EMG shows sparing of paraspinal muscles in a plexopathy because the lesion is, by definition, distal to the origin of the dorsal primary rami (cf. radiculopathy). Coexistence of radiculopathy and plexopathy may invalidate these simple rules.

- **Recognized causes of brachial plexopathy include**
  
  **Trauma:** Upper plexus: Dejerine–Klumpke paralysis (‘waiter’s tip’ posture)
  
  Lower plexus: Erb–Duchenne paralysis (claw hand)
  
  Inflammation/idiopathic: brachial neuritis, neuralgic amyotrophy
  
  Malignant infiltration, e.g. carcinoma of lung (Pancoast), breast, +/- Horner’s syndrome; pain a significant symptom
  
  Postradiation (e.g. after radiotherapy for malignant breast cancer with axillary spread; myokymic discharges may be seen on EMG)
  
  Tomaculous neuropathy
  
  Hereditary neuropathy with liability to pressure palsies (HNLPP)
  
  Neurogenic thoracic outlet syndrome (rare): cervical rib or C7 transverse process or fibrous band compressing the lower trunk; may be surgically remediable

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**Cross References**
Paresis; Weakness
Recognized causes of lumbosacral plexopathy include

- Compression; e.g. iliopsoas haematoma (anticoagulation, haemophilia), abscess (tuberculosis); abdominal aortic aneurysm; pregnancy (foetal head in the second stage of labour)
- Neoplasia (direct spread > metastasis)
- Trauma (rare; cf. brachial plexopathy)
- Postradiation
- Vasculitis (mononeuritis multiplex much commoner)
- Idiopathic

Imaging with MRI is superior to CT for defining structural causes of plexopathy.

**References**


**Cross References**

Amyotrophy; Claw hand; Horner’s syndrome; Nerve thickening; Neuropathy; Radiculopathy; ‘Waiter’s tip’ posture

**Polyganglionopathy**
- see NEUROPATHY

**Polymyoclonus**
- see MYOCLONUS

**Polyneuropathy**
- see NEUROPATHY

**Polyopia**

Polyopia, or polyopsia, or multiplication of images, is a visual illusory phenomenon in which a single target is seen as multiple images, most usually double but sometimes higher multiples (e.g. entomopia), persisting when looking away from the object. This may be likened to ‘echoes’ of the image, and eye movement may produce a trailing effect. Polyopia may be related to palinopsia.

Polyopia may occur as part of the visual aura of migraine and has also been associated with occipital and occipito-parietal lesions, bilateral or confined to the non-dominant hemisphere, and with drug abuse. It has also been described in disease of the retina and optic nerve and occasionally in normal individuals.

The pathophysiology is unknown; suggestions include a defect of visual fixation or of visual integration; the latter may reflect pure occipital cortical dysfunction.

**Reference**

Cross References
Entomopia; Illusion; Palinopsia

Popeye Arms
In facioscapulohumeral (FSH) muscular dystrophy, the deltoid muscle is normally well preserved, whilst biceps and triceps are weak and wasted, giving rise to an appearance of the upper limbs sometimes labelled as ‘Popeye arms’ or ‘chicken wings’.

Cross Reference
Winging of the scapula

Poriomania
A name sometimes given to prolonged wandering as an epileptic automatism, or a fugue state of non-convulsive status epilepticus.

Reference

Cross References
Automatism; Seizures

Porropsia
Porropsia, or teliopsia, is a form of metamorphopsia characterized by the misperception of objects as farther away from the observer than they really are (cf. pelopsia).

Cross References
Metamorphopsia; Pelopsia

Positional Manoeuvres
- see HALLPIKE MANOEUVR; HALLPIKE TEST; HEAD IMPULSE TEST; VESTIBULO-OCULAR REFLEXES

Posttetanic Potentiation
- see FACILITATION

Postural Hypotension
- see ORTHOSTATIC HYPOTENSION

Postural Reflexes
Postures such as standing are largely reflex in origin, dependent upon involuntary muscle contraction in antigravity muscles. Interference with such reflex activity impairs normal standing. Postural and righting reflexes depend not only on the integration of labyrinthine, proprioceptive, exteroceptive, and visual stimuli, mostly in the brainstem but also involve the cerebral cortex. However, abnormalities in these reflexes are of relatively little diagnostic value except in infants.

One exception is extrapyramidal disease (parkinsonism, Huntington’s disease, but not idiopathic dystonia) in which impairment or loss of postural reflexes may be observed. In the ‘pull test’ the examiner stands behind the patient, who
is standing comfortably, and pulls briskly on the shoulders; if balance is normal, the patient takes a step back; with impaired postural reflexes, this may provoke repetitive steps backwards (retropulsion, festination) or even en bloc falling, due to the failure of reflex muscle contraction necessary to maintain equilibrium. Pushing the patient forward may likewise provoke propulsion or festination, but this manoeuvre is less safe since the examiner will not be placed to catch the patient should they begin to topple over.

Cross References
Dystonia; Festinant gait, Festination; Parkinsonism; Proprioception; ‘Rocket sign’

Pourfour du Petit Syndrome
Pourfour du Petit syndrome is characterized by mydriasis, widening of the palpebral fissure, exophthalmos, hyperhidrosis (i.e. inverse Horner’s syndrome, sympathetic overactivity), flushing, and increased intraocular pressure due to irritation of the sympathetic chain in the neck.

Cross Reference
Horner’s syndrome

Pouting, Pout Reflex
The pout reflex consists of a pouting movement of the lips elicited by lightly tapping orbicularis oris with a finger or tendon hammer, or by tapping a spatula placed over the lips. This myotatic stretch reflex is indicative of a bilateral upper motor neurone lesion, which may be due to cerebrovascular small vessel disease, motor neurone disease or multiple sclerosis. It differs from the snout reflex, which refers to the reflex elicited by constant pressure on the philtrum. Hence the pout reflex is a phasic, the snout a tonic, response.

Reference

Cross References
Frontal release signs; Primitive reflexes

Prayer Sign
An inability to fully oppose the palmar surfaces of the digits with the hands held in the praying position, recognized causes of which include ulnar neuropathy (main en griffe), Dupuytren’s contracture, diabetic cheiroarthropathy, and camptodactyly.

Cross References
Camptodactyly; ‘Table top’ sign

Prehensile Thumb Sign
- see FROMENT’S SIGN

Presbyastasis
Presbyastasis, or the disequilibrium of ageing, is a condition of elderly patients who present with imbalance and disequilibrium that cannot be ascribed to a particular disease state or single causative factor (e.g. vestibular disease, visual impairment, peripheral neuropathy). It is thought that abnormal sensory input, abnormal CNS sensory processing, abnormal control mechanisms for balance,
Presbycusis
Presbycusis is a progressive sensorineural hearing loss, especially for high frequencies, developing with increasing age, which may reduce speech discrimination. It is thought to be due to age-related attrition of hair cells in the organ of Corti and/or spiral ganglion neurones.

Reference

Presbyopia
Presbyopia is progressive far-sightedness which is increasingly common with increasing age, thought to be due to an age-related impairment of accommodation.

Reference
Age-related signs

Prèsque vu
- see DÉJÀ VU

Pressure Provocation Test
This is one of the provocative tests for carpal tunnel syndrome: it is positive if paraesthesia in the distribution of the median nerve develop when pressure is exerted on the palmar aspect of the patient’s wrist at the level of the carpal tunnel for 60 s.

References
D’Arcy CA, McGee S. Does this patient have carpal tunnel syndrome? JAMA 2000; 283: 3110–3117.

Cross References
Flick sign; Phalen’s sign; Tinel’s sign

Prevost’s Sign
Also known as Vulpian’s sign, this refers to the acute and transient gaze palsy in a frontal lesion (e.g. infarct) which is towards the side of the lesion and away from the concurrent hemiparesis. The eyes can be brought to the other side with the oculocephalic manoeuvre or caloric testing. In contrast, thalamic and basal ganglia haemorrhages produce forced deviation of the eyes to the side contralateral to the lesion (wrong-way eyes).
**Priapism**

Priapism is an unintended, sustained, and usually painful erection of the penis unrelated to sexual activity. It may occur with intramedullary spinal cord lesions (e.g. multiple sclerosis) which damage the lumbosacral erection centres and has also been reported with lumbar canal spinal stenosis. There are also non-neurological causes, such as haematological conditions (sickle cell anaemia, polycythaemia rubra vera) which may cause intrapenile thromboses.

**Primitive Reflexes**

Reflexes which are normally found in infancy but which disappear with brain maturation during childhood may be labelled as ‘primitive reflexes’ if they re-emerge in adulthood as a consequence of pathological states. Many of these reflexes are seen with frontal lobe pathology (e.g. grasp, pout/snout, palmo-mental, rooting, corneomandibular) and hence may also be known as ‘frontal release signs’. However, the term ‘primitive reflex’ could equally apply to Babinski’s sign which is not necessarily frontal in origin.

**References**


**Cross References**

Babinski’s sign (1); Corneomandibular reflex; Frontal release signs; Grasp reflex; Palmo-mental reflex; Pout reflex; Rooting reflex

**Procerus Sign**

A focal dystonia of the procerus muscle, denoted the procerus sign, has been suggested to contribute to the ‘astonished’, ‘worried’, or ‘reptile-like’ facial expression typical of progressive supranuclear palsy, which may also be characterized by reduced blinking, lid retraction, and gaze palsy. All contrast with the hypomimia of Parkinson’s disease. It has also been described in corticobasal degeneration.

**References**


**Cross References**

Blinking; Dystonia; Hypomimia; Parkinsonism

**Pronator Drift**

Pronator drift is pronation of the forearm observed when the arms are held straightforward, palms up, with the eyes closed. It suggests a contralateral corticospinal tract lesion and may be accompanied by downward drift of the arm and flexion of the fingers and/or elbow. It reflects the relative weakness of supinators vs. pronators in the arm with a pyramidal lesion, in addition to the relative weakness of extensors vs. flexors. It may be an early sign of corticospinal tract dysfunction.
Proprioception

Proprioception sensation, or joint position sense, is knowledge about one's position in space, originating from sensory receptors in skin, muscle, and viscera. Proprioceptive information is carried within the dorsal columns of the spinal cord (more reliably so than vibration sensation, though not necessarily exclusively). Lesions affecting this part of the cord, particularly in the cervical region (e.g. subacute combined degeneration of the cord due to vitamin B₁₂ deficiency, tabes dorsalis), lead to impairments of proprioception with sparing of spinothalamic sensations (pin-prick, temperature) producing a dissociated sensory loss. Impairment of proprioception leads to sensory ataxia which may manifest clinically with pseudoathetosis or pseudochoreoathetosis (also seen in useless hand of Oppenheim) and with a positive Romberg’s sign.

Reference


Cross References

Ataxia; Dissociated sensory loss; Myelopathy; Pseudoathetosis; Pseudochoreoathetosis; Rombergism, Romberg’s sign; Useless hand of Oppenheim; Vibration

Proptosis

Proptosis is forward displacement of the eyeball, an exaggerated degree of exophthalmos. There may be lower lid retraction. Proptosis may be assessed clinically by standing directly behind the patient and gradually tipping the head back, observing when the globe of the eyeball first comes into view; this is most useful for asymmetric proptosis. An exophthalmometer may be used to measure proptosis.

Once established, it is crucial to determine whether the proptosis is axial or non-axial. Axial proptosis reflects increased pressure within or transmitted through the cone of extraocular muscles (e.g. thyroid ophthalmopathy, cavernous sinus thrombosis), whereas non-axial proptosis suggests pressure from an orbital mass outside the cone of muscles (e.g. orbital lymphoma, pseudotumour, mucoccele). Pulsatile axial proptosis may occur in carotico-cavernous fistula, in which case there may be a bruit audible by auscultation over the eye. Venous angioma of the orbit may cause an intermittent proptosis associated with straining, bending, coughing, or blowing the nose.

Dedicated orbital CT or MRI, the latter with fat-suppression sequences and intravenous gadolinium contrast, may be required to detect intraorbital masses.

Middle cranial fossa tumours may cause pressure on the veins of the cavernous sinus with secondary intraorbital venous congestion causing a ‘false-localizing’ proptosis.

Cross References

Exophthalmos; ‘False-localizing signs’; Lid retraction
Prosopagnosia

Prosopagnosia is a form of visual agnosia characterized by an inability to recognize previously known human faces or equivalent stimuli (hence, a retrograde defect) and to learn new ones (anterograde defect). As with more pervasive visual agnosia, this may be

- **apperceptive**: due to faulty perceptual analysis of faces; or
- **associative**: a semantic defect in recognition.

Familiar individuals may be recognized by their voices or clothing or hair; hence, the defect may be one of visually triggered episodic memory. It is important to note that the defect is not limited solely to faces; it may encompass animals (‘zooagnosia’) or cars.

Prosopagnosia is often found in association with a visual field defect, most often a left superior quadrantanopia or even hemianopia, although for the diagnosis of prosopagnosia to be made this should not be sufficient to produce a perceptual deficit. Alexia and achromatopsia may also be present, depending on the exact extent of the underlying lesion.

Anatomically, prosopagnosia occurs most often in association with bilateral occipito-temporal lesions involving the inferior and mesial visual association cortices in the lingual and fusiform gyri, sometimes with subjacent white matter. Unilateral non-dominant (right) hemisphere lesions have occasionally been associated with prosopagnosia, and a syndrome of progressive prosopagnosia associated with selective focal atrophy of the right temporal lobe has been reported. Involvement of the periventricular region on the left side may explain accompanying alexia, and disconnection of the inferior visual association cortex (area V4) may explain achromatopsia.

Pathological causes of prosopagnosia include

- Cerebrovascular disease: by far the most common cause;
- Tumour, e.g. glioma, extending from one hemisphere to the other via the splenium of the corpus callosum;
- Epilepsy (paroxysmal prosopagnosia), due to bilateral foci or spread from one occipital focus to the contralateral hemisphere;
- Focal right temporal lobe atrophy;
- Herpes simplex encephalitis, usually as part of an extensive amnesic syndrome (although memory impairment may put this outwith the operational criteria for an agnosia);
- Developmental (or ‘congenital’) prosopagnosia; suggests that facial recognition is a separate neuropsychological function (the acquired pathologies do not respect functional boundaries).

References


**Cross References**

Achromatopsia; Agnosia; Alexia; Hemianopia; Phonagnosia; Quadrantanopia; Visual agnosia; Zooagnosia

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**Prosopoplegia**

- see BELL’S PALSY; FACIAL PARESIS, FACIAL WEAKNESS

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**Proust Phenomenon**

The Proust phenomenon, named after the author Marcel Proust (1871–1922), is the observation that particular odours may trigger reminders of autobiographical memories. There is some experimental evidence that olfactory stimuli can cue autobiographical memories more effectively than cues from other sensory modalities. The ‘petite madeleines phenomenon’ has been used to describe sudden triggering of memories in individuals with amnesia due to thalamic infarction.

**References**


**Cross Reference**

Amnesia

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**Proximal Limb Weakness**

Weakness affecting predominantly the proximal musculature (shoulder abductors and hip flexors) is a pattern frequently observed in myopathic and dystrophic muscle disorders and neuromuscular junction transmission disorders, much more so than predominantly distal weakness (the differential diagnosis of which encompasses myotonic dystrophy, distal myopathy of Miyoshi type, desmin myopathy, and, rarely, myasthenia gravis). Some neuropathic disorders may also cause a predominantly proximal weakness (e.g. Guillain–Barré syndrome). Age of onset and other clinical features may help to narrow the differential diagnosis: painful muscles may suggest an inflammatory cause (polymyositis, dermatomyositis); fatiguability may suggest myasthenia gravis (although lesser degrees of fatigue may be seen in myopathic disorders); weakness elsewhere may suggest a specific diagnosis (e.g. face in facioscapulohumeral muscular dystrophy, diaphragm in acid maltase deficiency) and cachexia points to underlying malignant disease; calf pseudohypertrophy suggests Duchenne or Becker muscular dystrophy; autonomic features and posttetanic potentiation of reflexes occur in Lambert–Eaton myasthenic syndrome. Investigations (blood creatine kinase, neurophysiology, and muscle biopsy) may be required to determine exact diagnosis. Differential diagnosis includes
• **Myopathies:**
  Inflammatory: polymyositis, dermatomyositis;
  Progressive muscular dystrophies: Duchenne, Becker, limb-girdle, facioscapulohumeral (FSH);
  Metabolic: acid maltase deficiency; thyroid dysfunction, Cushing’s syndrome;
  Non-metastatic feature of malignant disease;
  Drug-induced: alcohol, steroids.

• **Neuromuscular junction transmission disorders:**
  Myasthenia gravis;
  Lambert–Eaton myasthenic syndrome.

• **Neuropathy:**
  Guillain–Barré syndrome.

**Cross References**
Facilitation; Fatigue

**Pseudoabducens Palsy**
- see ABDUCENS (VI) NERVE PALSY

**Pseudoachromatopsia**
Pseudoachromatopsia is failure on tests of colour vision (e.g. pseudoisochromatic plates) which is not due to central or peripheral achromatopsia, for example, due to visual neglect.

**Cross References**
Achromatopsia; Neglect

**Pseudoagnosia**
- see AGNOSIA

**Pseudo-Argyll Robertson Pupil**
A pseudo-Argyll Robertson pupil shows light-near dissociation of pupillary reactions, but unlike the ‘true’ Argyll Robertson pupil, there is no miosis or pupil irregularity. Indeed the pupil may be dilated (mydriasis) and resemble a Holmes–Adie pupil. The latter may be differentiated on the basis of its response to dilute (0.2%) pilocarpine: Holmes–Adie pupil results from a peripheral lesion and shows denervation supersensitivity, constricting with dilute pilocarpine, whereas the pseudo-Argyll Robertson pupil results from a central lesion and does not respond. Pseudo-Argyll Robertson pupil has been reported in:

- Diabetes mellitus
- Multiple sclerosis
- Wernicke’s encephalopathy
- Neurosarcoidosis
- Tumour
- Haemorrhage
- Aberrant oculomotor (III) nerve regeneration
- Spinocerebellar ataxia type 1 (SCA1)
Pseudoathetosis

Pseudoathetosis is the name given to athetoid-like movements, most usually of the outstretched fingers (‘piano-playing fingers’) and hands, resulting from sensory ataxia (impaired proprioception); it is worse with the eyes closed. There may also be chorea-like movements, hence pseudochoreoathetosis. Causes include any interruption to the anatomical pathway mediating proprioception, most often lesions in the dorsal cervical cord (e.g. multiple sclerosis, subacute combined degeneration of the cord due to vitamin B₁₂ deficiency, or nitrous oxide overuse), but also lesions of the large (myelinated) peripheral nerve fibres and of the parietal lobe.

Cross References
Athetosis; Chorea, Choreoathetosis; Proprioception; Pseudochoreoathetosis

Pseudo-Babinski Sign

Pseudo-Babinski sign is the name given to dystonic extension of the great toe on stroking the sole of the foot, as when trying to elicit Babinski’s sign, with which this may be confused, although pseudo-Babinski responses persist for longer, and spontaneous extension of the toe, striatal toe, may also be present. Pseudo-Babinski signs may normalize after dopaminergic treatment in dopa-responsive dystonia.

Reference

Cross References
Babinski sign (1); Striatal toe

Pseudobitemporal Hemianopia
- see HEMIANOPIA

Pseudobulbar Affect
- see EMOTIONALISM, EMOTIONAL LABILITY; PATHOLOGICAL CRYING, PATHOLOGICAL LAUGHTER; PSEUDOBULBAR PALSY

Pseudobulbar Palsy

Pseudobulbar palsy, or spastic bulbar palsy, describes bilateral upper motor neurone lesions affecting fibres passing to the cranial nerve nuclei (cf. bulbar palsy). This leads to a variety of clinical features, including

- difficulty with speech: spastic dysarthria, dysphonia;
- difficulty with swallowing: dysphagia;
- brisk jaw jerk and pout reflex; there may be trismus;
- slow, spastic, tongue movements;
- gag reflex may be depressed or exaggerated.

There may be associated emotional lability, or pathological laughter and crying (‘pseudobulbar affect’), and a gait disorder with *marche à petit pas*. There are
otherwise few signs in the limbs, aside from brisk reflexes and upgoing plantar responses (Babinski’s sign).

Recognized causes of pseudobulbar palsy include

- Motor neurone disease (in which there may be coincident bulbar palsy);
- Multiple sclerosis;
- Bilateral internal capsule lacunar infarctions, widespread small vessel disease (Binswanger’s disease);
- Congenital childhood suprabulbar palsy (Worster–Drought syndrome; perisylvian syndrome).

Pseudobulbar affect may respond to serotonin-reuptake inhibitors.

Reference

Cross References
Babinski’s sign (1); Bulbar palsy; Dysarthria; Dysphagia; Dysphonia; Emotionalism, Emotional lability; Gag reflex; Jaw jerk; Marche à petit pas; Pathological crying, Pathological laughter; Trismus; Upper motor neurone (UMN) syndrome

Pseudochoreoathetosis
Pseudochoreoathetosis is the name given to choreoathetoid-type involuntary movements, including dystonic movements, which result from a loss or impairment of proprioception. These may be observed with lesions anywhere along the proprioceptive pathways, including parietal cortex, thalamus (there may be associated ataxic hemiparesis and hemihypoaesthesia), spinal cord, dorsal root ganglia (neuronopathy), and mononeuropathy.

References

Cross References
Ataxic hemiparesis; Chorea, Choreoathetosis; Dystonia; Proprioception; Pseudoathetosis; Useless hand of Oppenheim

Pseudodementia
Pseudodementia is a label given to cognitive impairments resulting from affective disorders, most commonly anxiety and depression; the terms ‘dementia syndrome of depression’ and ‘depression-related cognitive dysfunction’ have also been used. The pattern of cognitive deficits in individuals with depression most closely resembles that seen in so-called subcortical dementia, with bradyphrenia, attentional, and executive deficits. In addition there may be evident lack of effort and application, frequent ‘No’ or ‘don’t know’ answers, approximate answers (Ganser phenomenon, vorbereiden), and evidence of mood disturbance (tearfulness). Memory loss for recent and distant events may be equally severe
Pseudodiplopia

(cf. temporal gradient of memory loss in dementia, e.g. due to Alzheimer’s disease). A 22-item checklist to help differentiate pseudodementia from Alzheimer’s disease has been described, based on clinical history, behaviour, and mental status.

The recognition of pseudodementia is important since the deficits are often at least partially reversible with appropriate treatment with antidepressants. However, it should be borne in mind that depression is sometimes the presenting symptom of an underlying neurodegenerative dementing disorder such as Alzheimer’s disease. Psychomotor retardation in dementia syndromes may also be mistaken for depression. Longitudinal assessment may be required to differentiate between these diagnostic possibilities.

References

Cross References
Attention; Bradyphrenia; Dementia; Ganser phenomenon; Psychomotor retardation

Pseudodiplopia
- see PALINOPSIA

Pseudohallucination
The term pseudohallucination has been used in different ways. In the European psychopathological tradition, it may refer simply to vivid visual imagery, whereas in the American arena it may refer to hallucinations that are recognized for what they are, i.e. the patient has insight. Some patients with dementia with Lewy bodies certainly realize that their percepts do not correspond to external reality and similar experiences may occur with dopamine agonist treatment.

Reference

Cross Reference
Hallucination

Pseudohypertrophy
- see CALF HYPERTROPHY; MUSCLE HYPERTROPHY

Pseudo-Internuclear Ophthalmoplegia
Pseudo-internuclear ophthalmoplegia is a disorder of eye movements with impaired adduction in one eye and horizontal nystagmus in the abducting eye (i.e. signs as in an internuclear ophthalmoplegia) but without an intrinsic brainstem lesion. This sign may be seen in:

- Myasthenia gravis (a diagnosis which is always worthy of consideration in a patient with an ‘isolated INO’) due to extraocular muscle weakness
- Pseudo-One-and-a-Half Syndrome -

- Brainstem compression due to subdural haematoma with transtentorial herniation
- Cerebellar mass lesion
- Guillain–Barré syndrome, Miller Fisher syndrome
- Thyroid ophthalmopathy
- Orbital pseudotumour

The preservation of rapid saccades despite restriction of eye movements in myasthenia gravis may result from selective sparing of pale global muscle fibres which generate high-speed movements.

**References**

**Cross References**
Internuclear ophthalmoplegia (INO); One-and-a-half syndrome

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**Pseudomyotonia**
The term pseudomyotonia has been used in various ways:

- It may be used to describe the clinical appearance of myotonia (slow muscular relaxation after contraction) in the absence of myotonic discharges on electromyography. Pseudomyotonia is most commonly observed as the slow-relaxing or ‘hung-up’ tendon reflexes (Woltman’s sign) of hypothyroidism, although other causes are described.
- Pseudomyotonia has also been used to describe difficulty opening the hand in cervical osteoarthritis, although muscle relaxation is normal; finger flexion on attempted extension has been explained as due to aberrant axonal regeneration of the C7 root.
- The term pseudomyotonia has also been used to describe neuromyotonia and myokymia (as, for example, in Isaacs syndrome), to distinguish it from myotonia.

**References**

**Cross References**
Myotonia; Neuromyotonia; Woltman’s sign

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**Pseudo-One-and-a-Half Syndrome**
Pseudo-one-and-a-half syndrome is the eye movement disorder of one-and-a-half syndrome without a brainstem lesion. Myasthenia gravis and Guillain–Barré syndrome are recognized causes.
Pseudopapilloedema

Pseudopapilloedema is the name given to elevation of the optic disc that is not due to oedema (i.e. intracranial pressure is not raised). There may or may not be visible drusen (hyaline bodies). In distinction to oedematous disc swelling, the nerve fibre layer is not hazy and the underlying vessels are not obscured; however, spontaneous retinal venous pulsation is usually absent, and haemorrhages may be seen, so these are not reliable distinguishing features. Visual acuity is usually normal, but visual field defects (most commonly in the inferior nasal field) may be found.

Cross References
Disc swelling; Papilloedema; Retinal venous pulsation

Pseudoparesis
- see PARESIS; WEAKNESS

Pseudoptosis
Ptosis, drooping of the eyelid, may need to be differentiated from pseudoptosis or functional ptosis. This may result simply from a redundant tarsal skin fold, especially in older patients, or be a functional condition. Frontalis underactivity may be a clinical indicator of the latter diagnosis (cf. compensatory overactivity of frontalis with other causes of ptosis).

The term pseudoptosis has also been used in the context of hypotropia; when the non-hypotropic eye fixates, the upper lid follows the hypotropic eye and appears ptotic, disappearing when fixation is with the hypotropic eye.

References

Cross Reference
Ptosis

Pseudoradicular Syndrome
Thalamic lesions may sometimes cause contralateral sensory symptoms in an apparent radicular (e.g. C8) distribution. If associated with perioral sensory symptoms this may be known as the cheiro-oral syndrome.

Reference

Pseudo-Von Graefe’s Sign
Pseudo-Von Graefe’s sign is involuntary retraction or elevation of the upper eyelid (cf. Von Graefe’s sign), medial rotation of the eye, and pupillary constriction.
Ptosis

Ptosis, or blepharoptosis, is the name given to drooping of the eyelid. This may be due to mechanical causes such as aponeurosis dehiscence, or neurological disease, in which case it may be congenital or acquired, partial or complete, unilateral or bilateral, fixed or variable, isolated or accompanied by other signs, e.g. miosis in a Horner’s syndrome; diplopia in myasthenia gravis; mydriasis and downward and outward deviation of the eye in an oculomotor (III) nerve palsy. Ptosis may result from pathology in a variety of locations: brainstem disease involving the oculomotor (III) nerve; anywhere along the oculosympathetic autonomic pathway causing a Horner’s syndrome; or cortical disease (e.g. infarction) reflecting hemispheric control of the eyelid (probably bilaterally represented).

When considering the cause of ptosis, the differential diagnosis is broad. Recognized causes include

- **Congenital:**
  - Cranial nerve dysinnervation disorder
  - Congenital Horner’s syndrome
  - Oculomotor-trigeminal (or trigeminal-levator) synkinesis: Marcus Gunn jaw-winking phenomenon, or inverse Marcus Gunn phenomenon (ptosis on jaw opening)

Cross References
Lid retraction; Synkinesia, Synkinesis; Von Graefe’s sign

Psychic Akinesia
- see ATHYMHORMIA

Psychic Blindness
- see VISUAL AGNOSIA

Psychic Paralysis of Gaze
- see BALINT’S SYNDROME; OCULAR APRAXIA

Psychomotor Retardation
Psychomotor retardation is a slowness of thought (bradyphrenia) and movement (bradykinesia) seen in psychiatric disorders, particularly depression. It may be confused with the akinesia of parkinsonism and with states of abulia or catatonia. Psychomotor retardation may also be a feature of the ‘subcortical’ type of dementia or of impairments of arousal (obtundation).

Cross References
Abulia; Akinesia; Catatonia; Dementia; Obtundation; Parkinsonism

Psychomotor Signs
- see FRONTAL RELEASE SIGNS

Ptarmus
- see SNEEZING

Ptosis
Ptosis, or blepharoptosis, is the name given to drooping of the eye. This constellation of findings is said to be a lid-gaze synkinesis following aberrant axonal regeneration after an oculomotor (III) nerve palsy, usually of traumatic or chronic compressive rather than ischaemic origin.
• **Neurogenic:**
  Supranuclear lesion:
  Hemiparesis: due to cortical infarct; ptosis usually ipsilateral, incomplete
  Duane syndrome: ptosis on eye adduction, due to supranuclear levator inhibition; usually with family history
  Oculomotor (III) nerve:
  Hypertension, diabetes mellitus: ptosis often complete; in a superior divisional third nerve palsy partial ptosis is associated with superior rectus weakness only
  Compressive lesion (e.g. posterior communicating artery aneurysm): ptosis usually incomplete; ptosis may be present with subarachnoid haemorrhage
  Guillain–Barré syndrome
  Facial paresis

• **Neuromuscular junction:**
  Myasthenia gravis: ptosis variable, bilateral or unilateral
  Excessive botulinum toxin, e.g. given for treatment of blepharospasm

• **Myogenic:** ptosis usually bilateral:
  Mitochondrial disease (CPEO)
  Myotonic dystrophy
  Oculopharyngeal muscular dystrophy (OPMD)

• **Local, ophthalmological causes:**
  Age-related aponeurosis dehiscence, trauma, thyroid eye disease, lid inflammation (chalazion), lymphoma

  Pseudoptosis (*q.v.*), enters the differential diagnosis.

  Enhanced ptosis, worsening of ptosis on one side when the other eyelid is held elevated in a fixed position, may be demonstrated in myasthenia gravis and Lambert–Eaton myasthenic syndrome.

**References**

**Cross References**
Blepharospasm; Diplopia; Divisional palsy; Ewart phenomenon; Horner’s syndrome; Ice pack test; Jaw winking; Miosis; Mydriasis; Pseudoptosis; Pupil sparing; Synkinesia, Synkinesis

**Ptyalism**
- see SIALORRHOEA

**Pulfrich Phenomenon**
The Pulfrich phenomenon is the observation that a pendulum swinging from side to side appears to traverse a curved trajectory. This is a stereo-illusion resulting from latency disparities in the visual pathways, most commonly seen as a
Pupillary Reflexes

consequence of conduction slowing in a demyelinated optic nerve following unilateral optic neuritis. A tinted coloured lens in front of the good eye can alleviate the symptom (or induce it in the normally sighted).

References

Cross References
Phosphene; Relative afferent pupillary defect (RAPD)

Pull Test
- see POSTURAL REFLEXES

Punding
Punding is characterized by repetitive pointless behaviours, with a compulsive flavour to them, carried on for long periods of time to the exclusion of other activities (like writing a book). It is frequently related to previous occupation or hobbies but is seldom pleasurable. It occurs in Parkinson’s disease but the incidence is low (1.4% in one study). It is thought to be related to dopaminergic stimulation and may be associated with impulse control disorder such as pathological gambling and hypersexuality.

Reference

Cross References
Gambling; Hypersexuality

Pupillary Reflexes
Two pupillary reflexes are routinely examined in clinical practice:

- **Light reflex:**
  The eye is illuminated directly and the reaction (constriction) observed; the *consensual light reflex* is observed by illuminating the contralateral eye. In an eye with poor visual acuity, a *relative afferent pupillary defect* may be observed using the ‘swinging flashlight test’. The afferent pathway subserving the light reflex is optic nerve to thalamus, brainstem, and Edinger–Westphal nucleus, with the efferent limb (pupillomotor parasympathetic fibres) in the oculomotor (III) nerve. The contralateral (consensual) response results from fibres crossing the midline in the optic chiasm and in the posterior commissure at the level of the rostral brainstem. Paradoxical constriction of the pupil in darkness (Flynn phenomenon) has been described.

- **Accommodation reflex:**
  This is most conveniently examined by asking the patient to look into the distance, then focus on a near object (sufficiently close to necessitate
convergence of the visual axes) when pupil constriction should occur (accommodation–convergence synkinesis). The afferent pathways subserving this response are less certain than for the light reflex and may involve the occipital cortex, although the final (efferent) pathway via Edinger–Westphal nucleus and oculomotor nerve is common to both accommodation and light reflexes.

In comatose patients, fixed dilated pupils may be observed with central diencephalic herniation, whereas midbrain lesions produce fixed midposition pupils.

A dissociation between the light and accommodation reactions (light-near pupillary dissociation, q.v.) may be observed.

Reference

Cross References
Argyll Robertson pupil; Ciliospinal response; Cortical blindness; Flynn phenomenon; Light-near pupillary dissociation; Miosis; Mydriasis; Relative afferent pupillary defect (RAPD); Swinging flashlight sign

Pupil Sparing
Oculomotor (III) nerve lesions may be pupil sparing (normal response to light) or pupil-involving (mydriasis, loss of light reflex). The latter situation usually implies a ‘surgical’ cause of oculomotor palsy (e.g. posterior communicating artery aneurysm), especially if extraocular muscle function is relatively preserved. Pupil sparing suggests a ‘medical’ cause (e.g. diabetes mellitus, hypertension) especially if the palsy is otherwise complete (complete ptosis, eye deviated downwards and outwards). This disparity arises because pupilomotor fibres run on the outside of the oculomotor nerve and are relatively spared by ischaemia but are vulnerable to external compression. However, the distinction is not absolute; imaging for an aneurysm (by means of spiral CT, MRA, or catheter angiography) may be necessary if the clinical scenario leaves room for doubt.

Cross References
Oculomotor (III) nerve palsy; Ophthalmoparesis, Ophthalmoplegia; Ptosis; Pupillary reflexes

Pure Word Blindness
- see ALEXIA

Pure Word Deafness
Pure word deafness is a rare condition characterized by an inability to comprehend and discriminate spoken language, despite adequate hearing as measured by audiometry and with preserved spontaneous speech, reading, reading comprehension, and writing (i.e. no aphasia, alexia, agraphia). Lip reading may assist in the understanding of others who sometimes seem to the patient as though they are speaking in a foreign language. Patients can copy and write spontaneously, follow written commands, but cannot write to dictation. Word repetition tasks
are impaired. There may be associated amusia, depending on the precise location of cerebral damage.

Pure word deafness has been variously conceptualized as a form of auditory agnosia or a subcortical sensory aphasia.

Pure word deafness is most commonly associated with bilateral lesions of the temporal cortex or subcortical lesions whose anatomical effect is to damage the primary auditory cortex or isolate it (e.g. from Wernicke’s area) through lesions of the auditory radiation; unilateral lesions producing this syndrome have been reported. Very rarely pure word deafness has been associated with bilateral brainstem lesions at the level of the inferior colliculi.

References


Cross References
Agnosia; Amusia; Aphasia

Pursuit

Pursuit, or smooth pursuit, eye movements hold the image of a moving target on the fovea, or during linear self-motion, i.e. they stabilize the gaze. This is dependent upon vestibulo-ocular reflexes and visually mediated reflexes. Impaired pursuit may result from occipital lobe lesions, and may be abolished by bilateral lesions, and may coexist with some forms of congenital nystagmus.

References


Cross References
Nystagmus; Saccades; Saccadic intrusion, Saccadic pursuit

Pyramidal Decussation Syndrome

Pyramidal decussation syndrome is a rare crossed hemiplegia syndrome, with weakness of one arm and the contralateral leg without involvement of the face, due to a lesion within the pyramid below the decussation of corticospinal fibres destined for the arm but above that for fibres destined for the leg.

Pyramidal Signs, Pyramidal Weakness

- see HEMIPARESIS; UPPER MOTOR NEURONE (UMN) SYNDROME; WEAKNESS
Quadrantanopia
Quadrantanopia (quadrantanopsia), a defect in one quarter of the visual field, suggests an optic radiation lesion. Occipital lobe pathology is the most common cause of both inferior and superior quadrantanopias, although temporal lobe pathology disabling Meyer’s loop typically must be considered with a superior homonymous quadrantanopia (‘pie-in-the-sky’ defect). Parietal lobe lesions may produce inferior quadrant defects, usually accompanied by other localizing signs. Damage to extrastriate visual cortex (areas V2 and V3) has also been suggested to cause quadrantanopia; concurrent central achromatopsia favours this localization.

References

Cross References
Achromatopsia; Hemianopia; ‘Pie-in-the-sky’ defect

Quadriparesis, Quadriplegia
Quadriparesis or quadriplegia (tetraparesis, tetraplegia) refers to weakness, partial or total, respectively, of all four limbs which may be of upper motor neurone or, less commonly, lower motor neurone type (e.g. in Guillain–Barré syndrome).

- Lower motor neurone, and some acute upper motor neurone, pathologies produce a flaccid quadriparesis/quadriplegia with areflexia; urinary retention may be present.
- Upper motor neurone lesions, particularly if chronic, produce a spastic quadriparesis with hypertonia, sustained clonus, hyperreflexia, loss of abdominal and cremasteric reflexes, and bilateral Babinski’s sign. As with hemiplegia, upper motor neurone quadriplegia may result from lesions of the corticospinal pathways anywhere from motor cortex to cervical cord via the brainstem, but is most commonly seen with brainstem and upper cervical cord lesions. In such circumstances, respiration may be affected. There may also be enhanced flexion defence reflexes (‘flexor spasms’) which may develop over time into a fixed flexion deformity with secondary contractures (‘paraplegia in flexion’). Incomplete or high spinal cord lesions may evolve to ‘paraplegia in extension’.

Cross References
Hemiparesis; Paraplegia

Quadrupedalism
Quadrupedalism, walking on all fours, has been observed as part of a recessive cerebellar hypoplastic syndrome associated with cerebellar ataxia and learning disability. This may result from mutations in the carbonic anhydrase-related protein 8 and has also been linked to two other loci, VLDLR and a locus on chromosome 17p.

Reference
**Rabbit Syndrome**

The rabbit syndrome is a rest tremor of perioral and nasal muscles. It has been associated with both antipsychotic drug therapy and idiopathic Parkinson's disease and is therefore presumably related to dopamine deficiency. No specific investigations are required, but a drug history, including over the counter medication, is crucial. The condition may be confused with edentulous dyskinesia, if there is accompanying tremor of the jaw and/or lip, or with tardive dyskinesia. Drug-induced rabbit syndrome may remit with drug withdrawal but not always. Appropriate treatment of Parkinson's disease may also improve the involuntary movements. Anticholinergics may be tried.

**Reference**


**Cross Reference**

Parkinsonism

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**Raccoon Eyes**

‘Raccoon eyes’ refers to an appearance of bilateral periorbital ecchymosis, appearing 48–72 h after an anterior basal skull fracture.

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**Radiculopathy**

A radiculopathy is a disorder of nerve roots, causing pain in a radicular distribution, paraesthesia, sensory diminution or loss in the corresponding dermatome, and lower motor neurone type weakness with reflex diminution or loss in the corresponding myotome. Radiculopathies may be single or multiple (polyradiculopathy, e.g. cauda equina syndrome). There may be concurrent myelopathy, typically of extrinsic or extramedullary type. Most radiculopathies are in the lumbosacral region (60–90%), followed by the cervical region (5–30%). Electrophysiological studies may be helpful in distinguishing radiculopathy from a neuropathy or plexopathy: sensory nerve action potentials (SNAPs) are normal for intrathecal root lesions, and EMG shows involvement of paraspinal muscles. Recognized causes of radiculopathy include

- **Structural lesions:**
  - Compression: disc protrusion: cervical (especially C6, C7), lumbar (L5, S1) >>> thoracic; bony metastases; spondylolisthesis; fracture; infection;
  - Root avulsion, e.g. C5/C6, ‘waiter’s tip’ posture; C8/T1, claw hand +/- Horner’s syndrome.
Raynaud’s Phenomenon

Raynaud’s phenomenon consists of intermittent pallor or cyanosis, with or without suffusion and pain, of the fingers, toes, nose, ears, or jaw, in response to cold or stress. It may be observed by asking the patient to put their hands in cold water. Raynaud’s phenomenon may occur in Raynaud’s disease (idiopathic, primary) or Raynaud’s syndrome (secondary, symptomatic). Recognized causes include connective tissue disease, especially systemic sclerosis: cervical rib or thoracic outlet syndromes; vibration white finger; hypothyroidism; and uraemia. Associated symptoms should be sought to ascertain whether there is an underlying connective tissue disorder (e.g. rash, arthralgia, myalgia, calcium deposits in the skin, dysphagia). History of use of power tools should be sought (vibration white finger). The differential diagnosis includes causalgia. For Raynaud’s syndrome, the treatment is that of the underlying cause where possible. For Raynaud’s disease, and Raynaud’s syndrome where there is no effective treatment of the underlying cause, non-drug treatment encompasses life style adjustment to avoid precipitants and use of heated gloves. Drug therapy includes oral vasodilators (calcium channel blockers, ACE inhibitors), antioxidants (probucol), and prostacyclin analogues (bolus, infusions). Beta-blockers should be avoided.

Reference
Reduplicative Paramnesia

**Cross References**
Asynergia; Ataxia; Dysdiadochokinesia; Dysmetria; Hypotonia, Hypotonus; Macrographia.

**Recruitment**
Recruitment, or loudness recruitment, is the phenomenon of abnormally rapid growth of loudness with increase in sound intensity, which is encountered in patients with sensorineural (especially cochlear sensory) hearing loss. Thus patients have difficulty with sounds of low-to-moderate intensity (‘Speak up, doctor’) but intense sounds are uncomfortably loud (‘There’s no need to shout, doctor!’). Speech discrimination is relatively unimpaired in conductive hearing loss.

‘Recruitment’ may also be used to refer to pathological ‘spread’ of tendon reflexes, implying broadening of their receptive field.

**Cross Reference**
Reflexes

**Recurrent Utterances**
The recurrent utterances of global aphasia, sometimes known as verbal stereotypies, stereotyped aphasia, or monophasia, are reiterated words or syllables produced by patients with profound non-fluent aphasia (e.g. Broca’s original case, Leborgne, who could only repeat ‘tan, tan, tan’, by which name he was known). The poet Charles Baudelaire (1821–1867) may have been reduced to a similar state following a stroke.

**References**

**Red Ear Syndrome**
Irritation of the C3 nerve root may cause pain, burning, and redness of the pinna. This may also occur with temporomandibular joint dysfunction and thalamic lesions.

**Reference**

**Reduplicative Paramnesia**
Reduplicative paramnesia is a delusion in which patients believe familiar places, objects, individuals, or events to be duplicated. The syndrome is probably heterogeneous and bears some resemblance to the Capgras delusion as described by psychiatrists.

Reduplicative paramnesia is more commonly seen with right (non-dominant) hemisphere damage; frontal, temporal, and limbic system damage has been implicated. This may occur transiently as a consequence of cerebrovascular disease, following head trauma, or even after migraine attacks, or more
persistently in the context of neurodegenerative disorders such as Alzheimer’s disease.

Reference

Cross References
Capgras delusion; Delusion; Paramnesia

**Reflexes**
Reflex action – a sensory stimulus provoking an involuntary motor response – is a useful way of assessing the integrity of neurological function, since disease in the afferent (sensory) limb, synapse, or efferent (motor) limb of the reflex arc may lead to dysfunction, as may changes in inputs from higher centres.

Different types of reflex may be distinguished. Muscle tendon reflexes (myotactic reflexes) may be either tonic (in response to a static applied force: ‘stretch reflex’) or phasic (in response to a brief applied force, for example, a blow from a tendon hammer to the muscle tendon). The latter are of particular use in clinical work because of their localizing value (see Table). However, there are no reflexes between T2 and T12, and thus for localization one is dependent on sensory findings, or occasionally cutaneous (skin or superficial) reflexes, such as the abdominal reflexes.

<table>
<thead>
<tr>
<th>Reflex</th>
<th>Root value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Jaw jerk</td>
<td>Trigeminal (V) nerve</td>
</tr>
<tr>
<td>Supinator (brachioradialis, radial)</td>
<td>C5, C6</td>
</tr>
<tr>
<td>Biceps</td>
<td>C5, C6</td>
</tr>
<tr>
<td>Triceps</td>
<td>C7</td>
</tr>
<tr>
<td>Finger flexion (digital)</td>
<td>C8, T1</td>
</tr>
<tr>
<td>Abdominal</td>
<td>T7–T12</td>
</tr>
<tr>
<td>Cremasteric</td>
<td>L1, L2</td>
</tr>
<tr>
<td>Knee (Patellar)</td>
<td>L3, L4</td>
</tr>
<tr>
<td>Hamstring</td>
<td>L5, S1</td>
</tr>
<tr>
<td>Ankle (Achilles)</td>
<td>(L5) S1 (S2)</td>
</tr>
<tr>
<td>Bulbocavernosus</td>
<td>S2, S3, S4</td>
</tr>
<tr>
<td>Anal</td>
<td>S4, S5</td>
</tr>
</tbody>
</table>

Tendon reflex responses are usually graded on a five-point scale:
−: absent (areflexia; as in lower motor neurone syndromes, such as peripheral nerve or anterior horn cell disorders; or acute upper motor neurone syndromes, e.g. ‘spinal shock’);
+/−: present only with reinforcement (Jendrassik’s manoeuvre); hyporeflexia;
+: normal;
++: brisk normal;
+++: pathologically brisk (hyperreflexia, as in upper motor neurone syndromes).
Reflex ‘spread’, or ‘recruitment’, for example, a finger jerk when eliciting the supinator or biceps jerk, is suggestive of corticospinal pathway (upper motor neurone) pathology, producing an enlarged receptive field for the reflex response; concurrent disruption of the local reflex arc may result in inverted reflexes.

Reflex responses may vary according to the degree of patient relaxation or anxiety (precontraction). Moreover, there is interobserver variation in the assessment of tendon reflexes (as with all clinical signs): a biasing effect of prior knowledge upon reflex assessment has been recorded.

There is also a class or ‘primitive’, ‘developmental’, or ‘psychomotor’ signs, present in neonates but disappearing with maturity but which may re-emerge with ageing or cerebral (especially frontal lobe) disease, hence sometimes known as ‘frontal release signs’.

**References**

**Cross References**
Age-related signs; Areflexia; Crossed adductor reflex; Facilitation; Frontal release signs; Hyperreflexia; Hyporeflexia; Inverted reflexes; Jendrassik’s manoeuvre; Lower motor neurone (LMN) syndrome; Primitive reflexes; Pupillary reflexes; Upper motor neurone (UMN) syndrome; Woltman’s sign. See also specific (named) reflexes

**Relative Afferent Pupillary Defect (RAPD)**
An afferent pupillary defect (APD), or relative afferent pupillary defect (RAPD), is an abnormal pupillary response in which the normally equal direct and consensual pupillary reflexes are asymmetric, the direct response being less than the consensual. This may be particularly evident using the ‘swinging flashlight’ test, in which the two pupils are alternately illuminated every 2–3 s in a darkened room. Quickly moving the light to the diseased side may produce pupillary dilatation (Marcus Gunn pupil). Subjectively, patients may note that the light stimulus seems less bright in the affected eye.

RAPD suggests an asymmetric optic nerve pathology, such as optic neuritis or tumour, causing a conduction defect; indeed this is the most sensitive sign of optic nerve pathology. Although visual acuity may also be impaired in the affected eye, and the disc appears abnormal on fundoscopy, this is not necessarily the case. Since RAPD depends on asymmetry of optic nerve conduction, no defect may be observed if both optic nerves are affected.

RAPD has also been described with lesions of the retina, optic chiasm, optic tract (contralateral), brachium of the superior colliculus, and pretectal nucleus (in the latter two situations without visual impairment).

**References**

Cross References
Amblyopia; Marcus Gunn pupil, Marcus Gunn sign; Pupillary reflexes; Swinging flashlight sign

Religiosity
- see HYPERRELIGIOSITY

Remote Atrophy
- see ATROPHY; ‘FALSE-LOCALIZING SIGNS’

Retinal Venous Pulsation
Venous pulsation is evident in the normal retina when observed with an ophthalmoscope, particularly at the margin of the disc. It is sometimes difficult to see and may be more obvious in the recumbent position because of higher pressure within the retinal veins in that position.

Venous pulsation is expected to be lost when intracranial pressure rises above venous pressure. This may be a sensitive marker of raised intracranial pressure and an early sign of impending papilloedema. However, venous pulsation may also be absent in pseudopapilloedema and sometimes in normal individuals. Hence, the reliability of this sign has been questioned.

References

Cross References
Papilloedema; Pseudopapilloedema

Retinitis Pigmentosa
Retinitis pigmentosa, or tapetoretinal degeneration, is a generic name for inherited retinal degenerations characterized clinically by typical appearances on ophthalmoscopy, with peripheral pigmentation of ‘bone-spicule’ type, arteriolar attenuation, and eventually unmasking of choroidal vessels and optic atrophy. Despite the name, there is no inflammation; the pathogenetic mechanism may be apoptotic death of photoreceptors. This process may be asymptomatic in its early stages, but may later be a cause of nyctalopia (night blindness), and produce a midperipheral ring scotoma on visual field testing.

A variety of genetic causes of isolated retinitis pigmentosa have been partially characterized:

- **autosomal recessive**: linked to chromosome 1q;
- **X-linked**: Xp11, Xp21;
- **autosomal dominant**: 3q, 6p, 8.
At least some of these are related to mutations in the gene for the rod cell protein rhodopsin.

In most cases, patients with retinitis pigmentosa have no associated systemic or extraocular abnormalities, but there are a number of multisystem disorders in which it occurs,

- Abetalipoproteinaemia (Bassen–Kornzweig syndrome; HARP syndrome);
- Alström’s syndrome;
- Cockayne syndrome;
- Friedreich’s ataxia;
- Lawrence–Moon–Bardet–Biedl syndrome;
- Mitochondrial disorders (e.g. Kearns–Sayre syndrome and NARP);
- Neuronal ceroid lipofuscinosis;
- Peroxisomal disorders, Refsum’s disease;
- Usher’s disease.

References

Cross References
Nyctalopia; Optic atrophy; Scotoma

Retinopathy
Retinopathy is a pathological process affecting the retina, with changes observable on ophthalmoscopy; dilatation of the pupil aids observation of the peripheral retina. Common causes include

- Diabetes mellitus: various abnormalities may occur, in both insulin-dependent (IDDM) and non-insulin-dependent (NIDDM) patients. ‘Background’ diabetic retinopathy is manifest as microaneurysms, dot and blot haemorrhages, hard exudates, and diffuse retinal oedema, all of which may be asymptomatic. Oedema and hard exudates at the macula are common causes of visual impairment. Proliferative retinopathy is characterized by neovascularization of the disc due to retinal hypoxia, typically in IDDM, with the risk of vitreous haemorrhage, traction retinal detachment, and irreversible visual loss. Laser treatment of new vessels is the treatment of choice
- Hypertension: hypertensive retinopathy may cause arteriolar constriction, with the development of cotton–wool spots; and abnormal vascular permeability causing flame-shaped haemorrhages, retinal oedema, and hard exudates; around the fovea, the latter may produce a macular star. Optic disc swelling may be seen in malignant hypertension. Arteriosclerosis, thickening of vessel walls with prolonged hypertension, may cause changes at arteriovenous crossings (‘AV nipping’). Systemic hypertension is associated with an increased risk of branch retinal vein and central retinal artery occlusion
- Drug-induced, e.g. antimalarials (chloroquine); chlorpromazine
• Retinitis pigmentosa
• Serous retinopathy or chorioretinopathy: leakage of fluid into the subretinal space, causing unilateral sudden non-progressive visual loss
• Cancer-associated retinopathy: arteriolar narrowing, optic atrophy
• ‘Salt and pepper’ retinopathy of Kearns–Sayre syndrome (mitochondrial disorder)

An electroretinogram (ERG) may be helpful in confirming the presence of a retinopathic disorder.

Cross References
Maculopathy; Retinitis pigmentosa; Scotoma

**Retrocollis**

Retrocollis is an extended posture of the neck. Progressive supranuclear palsy (PSP; Steele–Richardson–Olszewski syndrome) is commonly associated with retrocollis (cf. antecollis in multiple system atrophy). Retrocollis may also be a feature of cervical dystonia (torticollis) and of kernicterus.

Cross References
Antecollis; Dystonia; Parkinsonism; Torticollis

**Retropulsion, Retropulsion Test**
- see FESTINANT GAIT, FESTINATION; POSTURAL REFLEXES

**Reverse Lhermitte’s Sign**
- see LHERMITTE’S SIGN

**Reverse Phalen’s Manoeuvre**
- see PHALEN’S SIGN

**Reverse Ptosis**
- see HORNER’S SYNDROME; PTOSIS

**Reverse Sensory Geste**
Whereas the sensory geste antagoniste transiently reverses the severity of dystonia, reverse sensory geste worsens it, for example, in cervical dystonia (torticollis).

Reference

Cross References
Dystonia; Geste antagoniste; Torticollis

**Reverse Straight Leg Raising**
- see FEMORAL STRETCH TEST

**Revilliod’s Sign**
Revilliod’s sign is an acquired inability to wink. This is a sign, possibly early, of corticobulbar disease.
**Riddoch’s Phenomenon**
Riddoch’s phenomenon is the dissociation of the perception of static and kinetic visual stimuli (statokinetic dissociation). This phenomenon does not have particular localizing value, since it may occur with both occipital and anterior visual pathway lesions.

**Reference**

**Cross References**
Akinetopsia; Visual agnosia

**Right–Left Disorientation**
Right–left disorientation is an inability to say whether a part of the body is on the right or left side or to use a named body part to command. This may occur in association with acalculia, agraphia, and finger agnosia, collectively known as the Gerstmann syndrome. Although all these features are dissociable, their concurrence indicates a posterior parietal dominant hemisphere lesion involving the angular and supramarginal gyri.

**Cross References**
Acalculia; Agraphia; Autotopagnosia; Finger agnosia; Gerstmann syndrome

**Rigidity**
Rigidity is an increased resistance to the passive movement of a joint which is constant throughout the range of joint displacement and not related to the speed of joint movement; resistance is present in both agonist and antagonist muscles. In these particulars, rigidity differs from spasticity. Rigidity also needs to be differentiated from stiffness. Rigidity may be described as:

- **consistent**: ‘leadpipe rigidity’; or
- **jerky**: ‘cogwheel rigidity’ or Negro’s sign, when a rhythmic fluctuation (i.e. tremor), like a ratchet or cogwheel, is superimposed on the background of sustained rigidity (NB cogwheeling, reflecting underlying tremor, may occur in the absence of rigidity, e.g. in essential tremor).

The presence of rigidity may be made more obvious by reinforcing manoeuvres (e.g. clenching and relaxing the contralateral fist, performing mental arithmetic), a finding variously known as activated rigidity, or Froment’s sign, or synkinesis (but note that both Froment’s sign and synkinesis have other meanings too). However, this may occur in some normal subjects; it is most helpful in the diagnosis of Parkinson’s disease if unilateral. Rigidity may also be demonstrated using Wartenberg’s swing test.

Rigidity is a feature of parkinsonism and may coexist with any of the other clinical features of extrapyramidal system disease, but particularly akinesia (akineti-rigid syndrome); both are associated with loss of dopamine projections from the substantia nigra to the putamen. Rigidity is a feature of pathology within the basal ganglia.

The pathophysiology of rigidity is thought to relate to overactivity of tonic stretch reflexes in the spinal cord due to excessive supraspinal drive to spinal cord...
α-motor neurones following loss of descending inhibition as a result of basal ganglia dysfunction. In other words, there is a change in the sensitivity of the spinal interneurones which control α-motor neurones due to defective supraspinal control. Hence rigidity is a positive or release symptom, reflecting the operation of intact suprasegmental centres. The physiological correlate of this is the increased EMG activity found in rigid muscles with increased 1A afferent fibre activity, suggesting maintained α–β linkage. In support of this, pyramidotomy has in the past been shown to produce some relief of rigidity.

Rigidity in Parkinson’s disease may be lessened by treatment with levodopa preparations. The techniques of modern stereotactic neurosurgery may also be helpful, particularly stimulation of the subthalamic nucleus, although both thalamotomy and pallidotomy may also have an effect.

The term rigidity may also be used to describe

• posturing associated with coma: decorticate or decerebrate, flexor and extensor posturing, respectively;
• a lack of mental flexibility, particularly evident in patients with frontal lobe dysfunction.

References

Cross References
Decerebrate rigidity; Decorticate rigidity; Froment’s sign; Frontal lobe syndromes; Parkinsonism; Stiffness; Synkinesia, Synkinesis; Tremor; Wartenberg’s swing test

Rindblindheit
- see CORTICAL BLINDNESS

Ring Scotoma
- see ANNULAR SCOTOMA; SCOTOMA

Rinne’s Test
Rinne’s test is one of the tuning fork tests (512 Hz fork preferred), which is used to define whether there is a conductive element to hearing loss. The patient is asked to compare the loudness of a vibrating tuning fork held at the external auditory meatus (air conduction; AC) with the loudness of the fork held against
the mastoid process (bone conduction; BC); masking of the other ear, for example, by rubbing the tragus, is advised. Normally air conduction is louder (AC > BC). If bone conduction sounds louder (BC > AC), then this is indicative of a conductive hearing loss. In sensorineural hearing loss, AC and BC are diminished to a similar extent, and air conduction remains louder (AC > BC).

**Reference**

**Cross References**
Schwabach test; Weber’s test

**Rising Sign**
- see BABINSKI’S TRUNK–THIGH TEST

**Risus Sardonicus**
Risus sardonicus (‘sardonic smile’) due to spasm of the facial musculature is a classic feature of the neuromuscular syndrome of tetanus, now exceptionally rarely seen in the developed nations. Risus sardonicus may also occur in the context of dystonia, more usually symptomatic (secondary) than idiopathic (primary) dystonia.

**Cross References**
Dystonia; Oromandibular dystonia; Spasm

**Robot Syndrome**
- see ATHYMHORMIA

**Rocket Sign**
The so-called rocket sign is a toppling backwards, after jumping to the feet from the sitting position, due to postural instability, seen in progressive supranuclear palsy (PSP) and ascribed to frontal lobe dysfunction. A history of falls due to postural instability in the first year after disease onset is one of the mandatory inclusion criteria for the diagnosis of PSP.

**Cross References**
Parkinsonism; ‘Wheelchair sign’

**Roger’s Sign**
Roger’s sign, or the numb chin syndrome, is an isolated neuropathy affecting the mental branch of the mandibular division of the trigeminal (V) nerve, causing pain, swelling, and numbness of the lower lip, chin, and mucous membrane inside the lip. This is usually a sign of metastatic spread of cancer to the jaw.

Hypoaesthesia involving the cheek, upper lip, upper incisors, and gingiva, due to involvement of the infraorbital portion of the maxillary division of the trigeminal nerve (‘numb cheek syndrome’), is also often an ominous sign, resulting from recurrence of squamous cell carcinoma of the face infiltrating the nerve.

**References**
Apparent malfunction of self-winding (Rolex) watches, which depend on movement of the arm, may occur when they are worn on a hypokinetic, rigid arm; this may be the first sign of a parkinsonian syndrome.

Cross References
Parkinsonism; Wartenberg’s swing test

Rombergism, Romberg’s Sign
Romberg’s sign is adjudged present (or positive) when there is a dramatic increase in unsteadiness, sometimes with falls, after eye closure in a patient standing comfortably (static Romberg’s test). Before asking the patient to close his or her eyes, it is advisable to position one’s arms in such a way as to be able to catch the patient should they begin to fall. Patients may fall forward immediately on eye closure (‘sink sign’). These phenomena result from sensory ataxia (i.e. loss of proprioception from the feet), which occurs most commonly with posterior column spinal cord disease: Romberg’s sign may be seen in tabes dorsalis.

A modest increase in sway on closing the eyes may be seen in normal subjects and patients with cerebellar ataxia, frontal lobe ataxia, and vestibular disorders (towards the side of the involved ear); on occasion these too may produce an increase in sway sufficient to cause falls. Hence, Romberg’s test is not specific. Posturography is an attempt to quantify the Romberg test.

Large amplitude sway without falling, due to the patient clutching hold of the physician, has been labelled ‘psychogenic Romberg’s sign’, an indicator of functional stance impairment.

Heel–toe (tandem) walking along a straight line is sometimes known as the dynamic Romberg’s test.

References

Cross References
Ataxia; Functional weakness and sensory disturbance; Proprioception; Tandem walking

Roos Test
Roos test, or the elevated arm stress test, may be helpful in the diagnosis of vascular thoracic outlet syndrome, along with Adson’s test. The arm is held above the head with the elbow extended and the hand exercised. Development of numbness, pain, and paraesthesia, along with pallor of the hand, supports the diagnosis of thoracic outlet syndrome.
Ross’s Syndrome

Cross Reference
Adson’s test

Rooting Reflex
The rooting reflex is a turning of the head towards a tactile stimulus on the face or an object approaching the mouth, a normal response in infants which is lost during childhood. Its presence in adults is indicative of diffuse premotor frontal disease, this being a primitive reflex or frontal release sign.

Reference

Cross References
Age-related signs; Frontal release signs; Primitive reflexes

Rosenbach’s Sign
- see ABDOMINAL REFLEXES

Ross’s Syndrome
- see HOLMES–ADIE PUPIL, HOLMES–ADIE
Saccades
Saccades are rapid, ballistic, yoked movements of the eyes which bring the gaze to a new location in visual space. These movements may be performed voluntarily (tested clinically by asking the patient to ‘Look to your left, keeping your head still’, etc.) or reflexively, i.e. in response to an object of potential interest within the visual field (tested clinically by asking the patient to shift gaze from one of examiner’s hands to another). Internuclear ophthalmoplegia may be revealed when testing saccadic eye movements.

A number of parameters may be observed, including latency of saccade onset, saccadic amplitude, and saccadic velocity. An antisaccadic task (i.e. suppression of saccades to a novel visual stimulus) may be used to assess ease of saccade suppression. Of these, saccadic velocity is the most important in terms of localization value, since it depends on burst neurones in the brainstem (paramedian pontine reticular formation for horizontal saccades, rostral interstitial nucleus of the medial longitudinal fasciculus for vertical saccades). Latency involves cortical and basal ganglia circuits; antisaccades involve frontal lobe structures; and amplitude involves basal ganglia and cerebellar circuits (saccadic hypometria, with a subsequent correctional saccade, may be seen in extrapyramidal disorders such as Parkinson’s disease; saccadic hypermetria or overshoot may be seen in cerebellar disorders). Difficulty in initiating saccades may be described as ocular (motor) apraxia. Antisaccades may be poorly suppressed in Huntington’s disease. In Alzheimer’s disease, patients may make reflex saccades towards a target in an antisaccadic task (visual grasp reflex).

Assessment of saccadic velocity may be of particular diagnostic use in parkinsonian syndromes. In progressive supranuclear palsy slowing of vertical saccades is an early sign (suggesting brainstem involvement; horizontal saccades may be affected later), whereas vertical saccades are affected late (if at all) in corticobasal degeneration, in which condition increased saccade latency is the more typical finding, perhaps reflective of cortical involvement.

References

Cross References
Internuclear ophthalmoplegia (INO); Ocular apraxia; Ocular flutter; Opsoclonus; Parkinsonism; Saccadic intrusion, Saccadic pursuit; Square wave jerks
Saccadic Intrusion, Saccadic Pursuit
Saccadic intrusions are inappropriate saccades which interfere with visual fixation (static, or during motor pursuit: saccadic pursuit). Several types of saccadic intrusion are described, including ocular flutter, opsoclonus, and square wave jerks. Saccadic (cogwheel) pursuit is normal in infants and may be a non-specific finding in adults; however, it may be seen in Huntington's disease.

Cross References
Ocular flutter; Opsoclonus; Pursuit; Saccades; Saccadic intrusion, Saccadic pursuit; Square wave jerks

Saccadomania
- see OPSOCLONUS

Sacral Sparing
Sacral sparing is the preservation of pain and temperature sensation in sacral dermatomes when there is loss in the legs and trunk. This is a late, unusual, but diagnostic feature of a spinal cord lesion, usually an intrinsic (intramedullary) lesion but sometimes an extramedullary compression. Spastic paraparesis below the level of the lesion due to corticospinal tract involvement is invariably present by this stage of sacral sparing.

Sacral sparing is explained by the lamination of fibres within the spinothalamic tract: ventrolateral fibres (of sacral origin), the most external fibres, are involved later than the dorsomedial fibres (of cervical and thoracic origin) by an expanding central intramedullary lesion (e.g. glioma, ependymoma, syringomyelia).

Although sacral sparing is rare, sacral sensation should always be checked in any patient with a spastic paraparesis.

Cross References
Dissociated sensory loss; Myelopathy; Paraparesis

Saddle Anaesthesia
- see ANAESTHESIA; CAUDA EQUINA SYNDROME

Saturday Night Palsy
- see WRIST DROP

Savant Syndrome
Savant syndrome, previously idiot savant syndrome, refers to individuals with developmental disability yet displaying skills at a level inconsistent with their general intellectual functioning. The outstanding ability may be feats of memory (recalling names), calculation (especially calendar calculation), music, or artistic skills, often in the context of autism or pervasive developmental disorder.Obsolete classification of such abilities as superlative technical skill, hypermnnesia, calculating idiots, and calendar artists has been superseded by interest in how the disparities between these and general intellectual abilities come about and whether this is some form of 'release' phenomenon. Occasionally, skills
such as artistic ability may emerge in the context of neurodegenerative disease (Alzheimer’s disease, frontotemporal lobar degeneration).

**References**

**Scanning Speech**
Scanning speech is a motor speech disorder (i.e. a dysarthria) comprising slow, deliberate, dysprosodic, monotonic verbal output. It may be confused with non-fluent aphasia (Broca’s aphasia).

Scanning speech was originally considered a feature of cerebellar disease in multiple sclerosis (after Charcot), and the term is often used with this implication. However, cerebellar disease typically produces an ataxic dysarthria (variable intonation, interruption between syllables, ‘explosive’ speech) which is somewhat different from scanning speech. Scanning speech correlates with midbrain lesions, often after recovery from prolonged coma.

**Cross References**
Asynergia; Aphasia; Broca’s aphasia; Cerebellar syndromes; Dysarthria

**Scapula Alata**
- see WINGING OF THE SCAPULA

**Schizophrenia**
This term has been used to describe the language disorder in schizophrenia, which may be characterized by paraphasias and neologisms, loose connections between thoughts, tangential thinking, and delusional intrusions. The resulting output may be unintelligible and may resemble Wernicke’s aphasia.

**Cross References**
Delusion; Neologism; Paraphasia; Wernicke’s aphasia

**Schwabach Test**
In the Schwabach test, a vibrating tuning fork is held against the patient’s mastoid process, as in Rinne’s test, until it is no longer audible. The examiner then places the tuning fork over his/her own mastoid, hence comparing bone conduction with that of the patient. If still audible to the examiner (presumed to have normal hearing), a sensorineural hearing loss is suspected, whereas in conductive hearing loss the test is normal.

**Cross Reference**
Rinne’s test

**Scoliosis**
- see KYPHOSCOLIYSIS
Scotoma
A scotoma is a localized area of impaired vision within an otherwise normal visual field. Mapping of the defect may be performed manually, by confrontation testing, or using an automated system. In addition to the peripheral field, the central field should also be tested, with the target object moved around the fixation point. A central scotoma may be picked up in this way or a more complex defect such as a centrocaecal scotoma in which both the macula and the blind spot are involved. Infarction of the occipital pole will produce a central visual loss, as will optic nerve inflammation. Scotomata may be absolute (no perception of form or light) or relative (preservation of form, loss of colour). Blindsight may fall into the latter category. A scotoma may be physiological, as in the blind spot or angioscotoma, or pathological, reflecting disease anywhere along the visual pathway from retina and choroid to visual cortex. Various types of scotoma may be detected:

- Central scotoma;
- Caecocentral or centrocaecal scotoma;
- Arcuate scotoma;
- Annular or ring scotoma;
- Junctional scotoma;
- Junctional scotoma of Traquair;
- Peripapillary scotoma (enlarged blind spot).

Cross References
Altitudinal field defect; Angioscotoma; Blindsight; Blind spot; Central scotoma, Centrocaecal scotoma; Hemianopia; Junctional scotoma, Junctional scotoma of Traquair; Maculopathy; Papilloedema; Quadrantanopia; Retinitis pigmentosa; Retinopathy; Visual field defects

Scratch Test
The ‘scratch test’, or ‘direction of scratch’ test, examines perception of the direction (up or down) of a scratch applied to the anterior shin (for example, with the sharp margin of a paper clip). It has been claimed as a reliable test of posterior column function of the spinal cord. Errors in this test correlate with central conduction times and vibration perception threshold.

References

Cross References
Proprioception; Vibration

Seborrhoea
Seborrhoea is a greasiness of the skin which may occur in extrapyramidal disorders, particularly Parkinson’s disease.
Seizures

Cross Reference
Parkinsonism

Seelenblindheit
- see VISUAL AGNOSIA

Seizures
Seizures are sudden, paroxysmal episodes of neurological dysfunction with or without impairment of consciousness, which may be epileptic (i.e. due to abnormal synchronous electrical activity within the brain, either focally or generally) or non-epileptic in origin (‘pseudoseizures’, non-epileptic attack disorder). The two varieties may coexist. Seizure morphology may be helpful in establishing aetiology and/or focus of onset.

- **Epileptic:**
  - *Idiopathic generalized:* tonic–clonic (‘grand mal’); absence attack (‘petit mal’); myoclonic epilepsy.
  - *Partial:* simple (no impairment of consciousness), for example, jerking of one arm, which may spread sequentially to other body parts (Jacksonian march); or complex, in which there is impairment or loss of consciousness: may be associated with specific aura (olfactory, *déjà vu, jamais vu*) and/or automatisms (motor, e.g. cursive; or emotional, e.g. gelastic, dacrystic); limb posturing (salutatory, fencing posture) and pelvic thrusting may be seen in frontal lobe epilepsy. Secondary generalization of seizures of partial onset may occur.

  Investigation of partial seizures to exclude a symptomatic cause is recommended (MR imaging, EEG). Some are amenable to surgical intervention. Otherwise, as for idiopathic generalized epilepsies, various antiepileptic medications are available. Partial seizures may prove more resistant to treatment than generalized seizures.

- **Non-epileptic:**
  - Often long lasting, thrashing, pelvic thrusting, carpet burns, eyes closed and resist opening, tongue biting rare, may have incontinence; past history of physical or sexual abuse. Best treated with psychological approaches or drug treatment of underlying affective disorders; antiepileptic medications are best avoided.

  The differentiation of epileptic from non-epileptic seizures may be difficult; it is sometimes helpful to see a video recording of the attacks or to undertake in-patient video-telemetry.

References
Self-Mutilation
Self-injury to the point of mutilation, especially around the mouth, may be seen in certain neurological conditions, such as Lesch–Nyhan syndrome, Tourette syndrome, and neuroacanthocytosis.

Sensory Ataxia
- see ATAXIA; ROMBERGISM, ROMBERG’S SIGN

Sensory Geste
- see GESTE ANTAGONISTE

Sequential Paresis
Sequential, or ‘round the clock’, paresis or weakness refers to the sequential development of weakness in one arm, the ipsilateral leg, the contralateral leg, and contralateral arm (i.e. hemiparesis, triparesis, tetraparesis or quadriplegia). This pattern is highly suggestive of a foramen magnum lesion, usually a tumour but sometimes demyelination or other intrinsic inflammatory disorder, sequentially affecting the lamination of corticospinal fibres in the medullary pyramids.

Setting Sun Sign
The setting sun sign, or sunset sign, consists of tonic downward deviation of the eyes with retraction of the upper eyelids exposing the sclera. There may be down-beating nystagmus. Setting sun sign is a sign of dorsal midbrain compression in children with untreated hydrocephalus.

A similar appearance may also be observed in progressive supranuclear palsy (Steele–Richardson–Olszewski syndrome; Stellwag’s sign) and in Parinaud’s syndrome, but without the tonic downward deviation.

Cross References
Lid retraction; Nystagmus; Parinaud’s syndrome; Stellwag’s sign

Shadowing
A neurobehavioural disorder, occasionally seen in patients with dementia, in which the patient follows the spouse or carer around like a shadow.

Cross Reference
Dementia

Shin-Tapping
A modification of the heel–knee–shin test or heel–shin test in which the heel is tapped repetitively on the shin before sliding it down to the foot, claimed to be a better test of motor coordination.
Sialorrhoea
Sialorrhoea (drooling) is excessive salivation, possibly due to excess flow of saliva but more likely secondary to a reduced frequency of swallowing (e.g. in parkinsonian syndromes) or difficulty in swallowing (e.g. motor neurone disease, developmental perisylvian syndrome). Metallic poisonings (mercury, bismuth, lead) may also produce marked salivation (ptyalism).

If troublesome, treatment of sialorrhoea with anticholinergic agents may be tried (atropine, hyoscine), although they may cause confusion in Parkinson’s disease. In extreme cases, irradiation of the salivary glands has been used. Recently, the use of intraparotid injections of botulinum toxin has been found useful.

References

Sighing
Occasional deep involuntary sighs may occur in multiple system atrophy. Sighing is also a feature, along with yawning, of the early (diencephalic) stage of central herniation of the brainstem with an otherwise normal respiratory pattern. Sudden inspiratory or expiratory sighs are said to be a feature of the hyperkinetic choreiform dysarthria characteristically seen in choreiform disorders such as Huntington’s disease.

Reference

Signe de l’éventail (Fan Sign)
- see BABINSKI’S SIGN (1)

Signe de Rideau
Signe de rideau, or curtain sign, refers to the motion of the posterior pharyngeal wall towards the intact side, resembling the drawing of a curtain, in unilateral paresis of the superior pharyngeal constrictor muscle, as seen in unilateral vagus (X) nerve palsy.

Signe du Journal
- see FROMENT’S SIGN
Simian Hand
Simian hand or ape hand has been used to describe the atrophy of the thenar eminence with recession of the metacarpal bones of the thumb to the plane of the other metacarpal bones seen in median nerve lesions in the axilla or upper arm.

Cross Reference
Benediction hand

Simian Posture
- see PARKINSONISM

Simultanagnosia
Simultanagnosia is impaired perception of multi-element or multipart visual displays, such that pictures are described in a piecemeal manner. Recognition of single objects is preserved; this is likened to having a fragment or island of clear vision which may shift from region to region. Two types of simultanagnosia are described:

- **Dorsal:**
  An attentional limitation preventing more than one object being seen at a time; although superficially similar to apperceptive visual agnosia, with which it has sometimes been classified, patients with dorsal simultanagnosia can recognize objects quickly and accurately, but unattended objects are not seen. There may be inability to localize stimuli even when they are seen, manifest as visual disorientation. Reading is severely impaired. Patients may grope, as though blind. Dorsal simultanagnosia is associated with bilateral posterior parieto-occipital lesions and is one feature of Balint’s syndrome.

- **Ventral:**
  A limitation in the number of objects which can be recognized in unit time, i.e. there is no primary recognition problem in that individual shapes can be recognized. Ventral simultanagnosia is most evident during reading which is severely impaired and empirically this may be the same impairment as seen in pure alexia; otherwise deficits may not be evident, unlike dorsal simultanagnosia. Ventral simultanagnosia may be a form of associative visual agnosia. It is associated with left inferior temporo-occipital cortical lesions.

References

Cross References
Agnosia; Alexia; Balint’s syndrome; Visual agnosia; Visual disorientation

Singultus
- see HICCUPS
Sink Sign
- see ROMBERG’S SIGN, ROMBERGISM

Skew Deviation
Skew deviation, or the Hertwig–Magendie sign, is a supranuclear vertical misalignment of the visual axes; the final common efferent pathway for eye movements is spared (cf. hypertropia, hypotropia, due to ocular motor nerve palsies or extraocular muscle disease). This is thought to reflect damage to otolith-ocular pathways or vestibulo-ocular pathways. There may be concurrent ocular tilt reaction. Bielschowsky’s head tilt test is usually negative (cf. ocular motor nerve palsies).

Skew deviation has been associated with posterior fossa lesions, from midbrain to medulla. Ipsiversive skew deviation (ipsilateral eye lowermost) has been associated with caudal pontomedullary lesions, whereas contraversive skew (contralateral eye lowermost) occurs with rostral pontomesencephalic lesions, indicating that skew type has localizing value.

Reference

Cross References
Bielschowsky’s sign, Bielschowsky’s test; Hypertropia; Hypotropia; Ocular tilt reaction; Tullio phenomenon

Smile–Wink Phenomenon
This name has been given to narrowing of the palpebral fissure aggravated by smiling following a contralateral lenticulocapsular infarction. Dysarthria, facial paresis, hemiparesis with or without hemihypoaesthesia, and excessive laughing with or without crying were common accompanying features in one series.

Reference

Cross References
Dysarthria; Facial paresis; Hemiparesis; Hypoaesthesia

Smooth Pursuit
- see PURSUIT

Snarling Facies
- see ‘MYASTHENIC SNARL’

Sneezing
Sneezing, or ptarmus, or sternutation, is a complex respiratory reflex. Sensory nasal trigeminal afferents run to a putative sneeze centre, localized to the brainstem based on lesions causing loss of sneezing following lateral medullary syndrome and medullary neoplasm. Integration of inputs in this centre reaches a threshold at which point an expiratory phase occurs with exhalation, forced eye closure, and contraction of respiratory musculature.
Sneezing may also be triggered by the presence of light: photoptalmosis or the photic sneeze reflex.

**References**


**Cross Reference**

Lateral medullary syndrome

**Snoring**

Reduced muscle tone in the upper airway during sleep leads to increased resistance to the flow of air, and partial obstruction often results in loud snoring. This symptom may be associated with the obstructive sleep apnoea–hypopnoea syndrome (OSAHS), risk factors for which include obesity, alcohol overconsumption, and male sex. OSAHS may be associated with a variety of neurological symptoms including excessive daytime somnolence, episodic loss of consciousness, headache (especially morning), cognitive decline, and increased risk of stroke (snoring may be an independent risk factor for stroke).

**References**


**Cross Reference**

Hypersomnolence

**Snouting, Snout Reflex**

Sometimes used interchangeably with pout reflex, this term should probably be reserved for the puckering or pouting of the lips induced by constant pressure over the philtrum, rather than the phasic response to a tap over the muscle with finger or tendon hammer.

**Reference**


**Cross References**

Frontal release signs; Pout reflex; Primitive reflexes

**Somatoparaphrenia**

Ascription of hemiplegic limb(s) to another person (e.g. the examiner, a family member). This may be a form of asomatognosia, or possibly a confabulation.
Spastic Catch

Reference

Cross References
Anosognosia; Asomatognosia; Autotopagnosia; Confabulation

Somatotopagnosia
- see AUTOTOPAGNOSIA

Spasm
The word spasm implies a sudden, involuntary, muscle contraction, which may be painful (cramp). For example, flexor spasms in patients paraplegic due to upper motor neurone lesions are sudden contractions of the flexor musculature, particularly of the legs, either spontaneous or triggered by light touch. Hemifacial spasm is an involuntary contraction of facial musculature. Tonic spasms (paroxysmal dystonia) occur in multiple sclerosis.

Spasm may also refer to a tetanic muscle contraction (tetany), as seen in hypocalcaemic states (e.g. *main d'accoucheur*), tetanus (e.g. risus sardonicus), or tonic spasms of various muscles (e.g. jaw musculature, trismus) which may be dystonic or spastic in origin. Involuntary movements such as tics may be known as spasms or habit spasms.

Patients may use the word spasm differently, e.g. to denote paroxysmal sensory phenomena, or even epileptic seizures. Infantile seizures consisting of brief flexion of the trunk and limbs (emposthotonos, salaam or jack-knife seizures) may be known as spasms.

Reference

Cross References
Contracture; Dystonia; Hemifacial spasm; *Main d'accoucheur*; Paraplegia; Risus sardonicus; Seizures; Tic; Tonic spasms; Trismus

Spasmus Nutans
Spasmus nutans is the clinical triad of head nodding, anomalous head postures, and nystagmoid eye movements seen in children aged between 1 and 8 years. This is usually a benign idiopathic condition, but the diagnosis should prompt consideration of an optic pathway tumour.

Reference

Cross Reference
Nystagmus

Spastic Catch
- see SPASTICITY
Spasticity

Spasticity is an increased resistance to the passive movement of a joint due to abnormally high muscle tone (hypertonus) which varies with the amplitude and speed of displacement of a joint (cf. rigidity). The excessive resistance evident at the extremes of joint displacement may suddenly give way, a phenomenon known as clasp-knife (or, confusingly, clasp-knife rigidity). Spasticity may vary in degree from mild (e.g. a spastic catch on supination/pronation of the forearm) to extreme (e.g. immobile limbs in fixed flexion with secondary contractures and painful spasms: paraplegia in flexion). Spasticity may need to be differentiated clinically from rigidity and stiffness.

The amount and pattern of spasticity depends on the location of the lesion and tends to be greater with spinal cord than cortical lesions. Scales to quantify spasticity are available (Ashworth, modified Ashworth, pendulum test of Wartenberg) but have shortcomings. Spasticity may also vary in distribution: for lesions above the spinal cord it typically affects the arm flexors and the leg extensors to a greater extent (hemiparetic posture).

Spasticity is a clinical feature of the upper motor neurone syndrome and may be accompanied by both positive (clonus, hyperreflexia, Babinski’s sign, flexor, or extensor spasms) and negative phenomena (weakness in a pyramidal distribution, motor underactivity): the latter may be more significant determinants of disability. Slow, laboured speech, with slow voluntary tongue movements, may be referred to as spastic dysarthria, which may occur in the context of a pseudobulbar palsy.

The pathogenesis of spasticity has traditionally been ascribed to damage to the corticospinal and/or corticobulbar pathways at any level from cerebral cortex to spinal cord. However, various lines of evidence (e.g. the failure of pyramiidotomy to produce spasticity in animals, rare human cases of isolated pyramid infarction causing hyperreflexia and weakness without spasticity) have led to the implication of other motor tracts in the genesis of spasticity, viz.:

- the dorsal reticulospinal tract, which lies in the lateral funiculus of the spinal cord and hence is often damaged concurrently with the adjacent lateral corticospinal tract (e.g. in MS, which seems to have a predilection for the lateral funiculus); this descending pathway has an inhibitory effect on stretch reflexes which is under cortical control;
- the medial reticulospinal tract and vestibulospinal tracts which are not under cortical control and whose excitatory effects on extensor tone may remain unopposed.

Physiologically, spasticity has been characterized as an exaggeration of the muscle stretch reflexes, with reduced threshold (hyperexcitable α-motor neurones) and abnormal reflex transmission (increased gain). The role of neurotransmitters (glutamate, glycine, catecholamines, serotonin) in the pathogenesis of spasticity is unclear, but the efficacy of baclofen (a GABA<sub>B</sub> agonist) and benzodiazepines suggests that impaired GABAergic transmission may contribute, perhaps through a loss of presynaptic inhibition mediated by interneurones or the inhibition of glutamate release.

Treatment of severe spasticity, for example, in multiple sclerosis, often requires a multidisciplinary approach. Urinary infection, constipation, skin
ulceration, and pain can all exacerbate spasticity, as may inappropriate posture; appropriate management of these features may ameliorate spasticity. Drugs which may be useful include baclofen, dantrolene (a blocker of muscle excitation–contraction coupling), and tizanidine (\(\alpha_2\)-adrenoreceptor agonist). Intrathecal baclofen given via a pump may also be of benefit in selected cases, and for focal spasticity injections of botulinum toxin may be appropriate. For painful immobile spastic legs with reflex spasms and double incontinence, irreversible nerve injury with intrathecal phenol or alcohol may be advocated to relieve symptoms. The place of cannabinoids has yet to be fully determined.

References

Cross References
Babinski’s sign (1); Clasp-knife phenomenon; Clonus; Contracture; Dysarthria; Hyperreflexia; Hypertonus; Paraplegia; Pseudobulbar palsy; Reflexes; Spasm; Upper motor neurone (UMN) syndrome; Weakness

Speech Apraxia
Speech apraxia is one of the labels applied to a disorder of communication characterized by slow speech tempo (‘groping for words’), impaired articulation, and dysprosody, with relatively intact language function and no dysgraphia. More errors occur with increasing articulatory complexity (consonant clusters vs. single consonants). Automatic or reactive speech (e.g. expletives, cliches) is without error. This, or a very similar, constellation of features has also been known as cortical dysarthria, aphemia, or phonetic disintegration. There may be associated oro-facial apraxia.

Speech apraxia has been associated with inferior frontal dominant (left) hemisphere damage in the region of the lower motor cortex or frontal operculum; it has been claimed that involvement of the anterior insula is specific for speech apraxia.

The exact nosological status of this entity remains in some doubt. The syndrome is thought to reflect disturbances of planning articulatory and phonatory functions, but is most often encountered as part of a non-fluent aphasia.

References

Cross References
Aphasia; Aphemia; Apraxia

Spinal Mass Reflex
The spinal mass reflex is involuntary flexion of the trunk in a comatose patient, such that they appear to be attempting to sit up (‘rising from the dead’).
**Cross Reference**
Coma

**Spoonerisms**
This term is used for a speech production disorder characterized by the transposition of consonants, so named for the mannerism affecting the speech of Reverend WA Spooner, Warden of New College, Oxford (1844–1930). If not deliberate, it presumably reflects a left hemisphere dysfunction in the appropriate sequencing of phonemes.

**Reference**

**Spurling’s Sign**
This is the name given to increase in arm pain (brachalgia) associated with compressive cervical radiculopathy following neck rotation and flexion to the side of the pain. A variant of this foraminal compression test involves rotation, side bend, and slight extension of the neck with the application of axial pressure to the head.

**Cross Reference**
Radiculopathy

**Square Wave Jerks**
Square wave jerks are small saccades which interrupt fixation, moving the eye away from the primary position and then returning. This instability of ocular fixation is a disorder of saccadic eye movements in which there is a saccadic interval (of about 200 ms; cf. ocular flutter, opsoclonus). Very frequent square wave jerks may be termed square wave oscillations. Very obvious square wave jerks (amplitude > 7°) are termed macrosquare wave jerks.

Square wave jerks are often best appreciated on ophthalmoscopy. Their name derives from the appearance they produce on electrooculographic recordings.

Although square wave jerks may be normal in elderly individuals, they may be indicative of disease of the cerebellum or brainstem, e.g. Huntington’s disease, Parkinson’s disease, progressive supranuclear palsy, cerebellar degeneration including multiple system atrophy. They have been reported to have close to 100% sensitivity for the diagnosis of PSP.

**Reference**
Troost BT, Daroff RB. The ocular motor defects in PSP. *Annals of Neurology* 1977; **2**: 397–403.

**Cross References**
Nystagmus; Ocular flutter; Opsoclonus; Saccadic intrusion, Saccadic pursuit

**Square Wave Oscillations**
- see SQUARE WAVE JERKS

**Squint**
- see HETEROTROPIA
Stapedius Reflex
- see HYPERACUSIS

Stellwag’s Sign
Stellwag’s sign is a widening of the palpebral fissure due to upper eyelid retraction. Along with a reduced blink rate, this creates a very typical staring, ‘astonished’, facies. The clinical phenomena of Stellwag’s sign overlap with those labelled as the sunset sign. Stellwag’s sign is seen in progressive supranuclear palsy and in dysthyroid eye disease.

Cross References
Blinking; Lid lag; Lid retraction; Sunset sign

Steppage, Stepping Gait
Steppage or stepping gait occurs with a lower motor neurone type of foot drop (‘floppy’ foot drop), e.g. due to a common peroneal nerve palsy, or peripheral neuropathies. Because of the weakness of foot dorsiflexion (weak tibialis anterior) there is compensatory overaction of hip and knee flexors during the swing phase of walking to ensure the foot clears the ground (hence ‘high-stepping gait’).
In the strike phase, there is a characteristic slapping down of the foot, again a consequence of weak ankle dorsiflexion. Proprioceptive loss, as in dorsal column spinal disease, may also lead to a gait characterized by high lifting of the feet and also stomping (stamping with a heavily accented rhythm) or slapping of the foot onto the floor in the strike phase.

The pattern of gait with upper motor neurone foot drop (‘stiff’ foot drop), e.g. due to a corticospinal tract lesion, is quite different, with the foot being dragged, sometimes with circumduction of the leg. This may lead to falls as a consequence of tripping over the foot, especially on up-hill gradients, and a characteristic pattern of wear on the point of the shoe.

Cross References
Foot drop; Lower motor neurone (LMN) syndrome; Proprioception; Rombergism, Romberg’s sign; Upper motor neurone (UMN) syndrome

Stereoanaesthesia
- see ASTEROEIGNOSIS

Sterehypaesthesia
- see ASTEROEIGNOSIS

Stereotyped Aphasia
- see RECURRENT UTTERANCES

Stereotypy
Stereotypies, or adventitious movements, may be defined as regular repeated movements, which are voluntary but not apparently goal-directed, and which may be carried out in a uniform pattern for long periods of time (cf. tic). Whole areas of the body may be involved by stereotypies and hence this movement is more complex than a tic. Examples include patting, tapping, rubbing, clasping,
wringing, digit sucking, body or head rocking or banging, grimacing, smelling, licking, spitting, and mouthing of objects.

Stereotypies are common in patients with learning disability, autism, and schizophrenia. Very characteristic manual stereotypies (washing, rubbing movements: ‘hand washing’) may be seen in Rett’s disease. The term has also been used to describe movements associated with chronic neuroleptic use; indeed adult-onset stereotypy is highly suggestive of prior exposure to dopamine-receptor-blocking drugs.

The recurrent utterances of global aphasia are sometimes known as verbal stereotypies or stereotyped aphasia. Reiterated words or syllables are produced by patients with profound non-fluent aphasia (e.g. Broca’s original case, Leborgne, who could only repeat ‘tan, tan, tan’, by which name he was known).

References

Cross References
Aphasia; Broca’s aphasia; Recurrent utterances; Tic

Sternocleidomastoid Test
It has been reported that apparent weakness of the sternocleidomastoid muscle is common (80%) in functional hemiparesis, usually ipsilateral to the hemiparesis, whereas it is rare in vascular hemiparesis (11%), presumably because of the bilateral innervation of the muscle.

Reference

Cross References
Functional weakness and sensory disturbance; Hemiparesis

Stethoscope Loudness Imbalance Test
- see HYPERACUSIS

Stewart–Holmes Sign
- see REBOUND PHENOMENON

Stiffness
Stiffness of muscles occurs as a feature of all pyramidal and extrapyramidal disorders (as spasticity and rigidity, respectively), but the term stiffness is usually reserved for disorders in which stiffness is the principal symptom due to continuous motor unit activity within muscles. There may be associated muscle pain (cramp). Stiffness may be primarily of muscular origin (myotonia) or of neural origin (myokymia, neuromyotonia). Accompanying signs may prove...
helpful in diagnosis, such as slow muscle relaxation (myotonia), percussion irritability of muscle (myoedema), and spontaneous and exertional muscle spasms. Hyperlordotic posture is typical of stiff man/stiff person syndrome. Stiffness must be differentiated from both rigidity and spasticity. Recognized causes of stiffness include the following:

- Stiff man/stiff person syndrome;
- Stiff limb syndrome;
- Progressive encephalomyelitis with rigidity +/− myoclonus;
- Neuromyotonia (Isaac’s syndrome; armadillo syndrome; peripheral nerve hyperexcitability);
- Schwartz–Jampel syndrome (chondrodystrophic myotonia);
- Tetanus;
- Strychnine poisoning.

The stiff man/stiff person syndrome is probably of autoimmune pathogenesis since it is strongly associated with insulin-dependent diabetes mellitus and the presence of antibodies to glutamic acid decarboxylase (anti-GAD antibodies), the enzyme in the synthetic pathway of GABA. Intravenous immunoglobulin therapy may be of symptomatic benefit.

References

Cross References
Myokymia; Myotonia; Neuromyotonia; Paramyotonia; Rigidity; Spasticity

**Stork Legs**
A name given to describe the disproportionate wasting of the lower legs, a pattern characteristic of hereditary motor and sensory neuropathies (Charcot–Marie–Tooth diseases), which may be evident even before the development of gait disorder with foot drop and steppage gait.

Cross References
Foot drop; Steppage, Stepping gait; Wasting

**Stork Manoeuvre**
The patient is asked to stand on one leg, with arms folded across chest, and the eyes open. Absence of wobble or falling is said to exclude a significant disorder of balance or pyramidal lower limb weakness.

**Strabismus**
- see HETEROPHORIA; HETEROTROPIA
**Straight Leg Raising**
- see LASEGUE’S SIGN

**Straight Thumb Sign**
Median nerve lesions in the forearm cause weakness of flexor pollicis longus, which normally flexes the distal phalanx of the thumb. Hence the thumb remains straight when the patient attempts to grasp something or make a fist. The ‘pinch sign’ may also be present.

**Reference**

**Cross Reference**
‘Pinch sign’

**Striatal Toe**
Striatal toe refers to the spontaneous tonic extension of the hallux which is seen in dystonic syndromes, and as a feature of extrapyramidal disorders, such as dopa-responsive dystonia. Striatal toe may be confused with Babinski’s sign (extensor plantar response) and pseudo-Babinski’s sign (= ‘phasic striatal toe’), the principal difference being that both the latter are elicited by stimulation whereas the former is a tonic response.

**Reference**

**Cross References**
Babinski’s sign (1); Parkinsonism; Pseudo-Babinski’s sign

**String Sign**
The string sign has been advocated as a way of testing visual field integrity in patients whose cooperation cannot be easily gained, by asking them to point quickly to the centre of a piece of string held horizontally in the examiner’s hands. If visual fields are full, the patient will point to the approximate centre; if there is a left field defect, pointing will be to the right of centre, and vice versa for a right field defect. Altitudinal field defects may be similarly identified by holding the string vertically.

**Reference**

**Cross Reference**
Visual field defects

**Stupor**
Stupor is a state of altered consciousness characterized by deep sleep or unresponsiveness, in which patients are susceptible to arousal only by vigorous and/or repeated stimuli, with lapse back into unresponsiveness when the stimulus stops. Stupor is a less severe impairment of conscious level than coma, but worse than obtundation (torpor). It is suggestive of diffuse cerebral dysfunction, e.g. drug-induced.
Sundowning

References

Cross References
Coma; Delirium; Encephalopathy; Obtundation

Stutter
Stutter, one of the reiterative speech disorders, is usually a developmental problem, but may be acquired in aphasia with unilateral or bilateral hemisphere lesions (e.g. vascular damage, trauma, Alzheimer’s disease, Parkinson’s disease, progressive supranuclear palsy). Unlike developmental stutter, acquired stutter may be evident throughout sentences, rather than just at the beginning. Furthermore, developmental stutter tends to occur more with plosives (phonemes where the flow of air is temporarily blocked and suddenly released, as in ‘p’, ‘b’), whereas acquired stutter is said to affect all speech sounds fairly equally. Cessation of developmental stutter following bilateral thalamic infarction in adult life has been reported, as has onset of stutter after anterior corpus callosum infarct.

References

Cross References
Aphasia; Echolalia; Palilalia

Sucking Reflex
Contact of an object with the lips will evoke sucking movements in an infant. The reflex may re-emerge in dementia.

Cross References
Akinetic mutism; Dementia; Frontal release signs

Summerskill’s Sign
- see LID RETRACTION

Sundowning
‘Sundowning’, or sundown syndrome, is increased confusion, agitation, or disorientation in the late afternoon, evening, and night-time, which may be seen in patients with delirium, and sometimes in dementia. In dementia, there may be complete reversal of sleep schedule with daytime somnolence and nocturnal wakefulness.

Although this syndrome may relate to worsening of visual cues with increasing darkness, it may also occur in well-lit environments. A disorder of circadian rhythms is a possible physiological correlate of ‘sundowning’: EEG recordings in delirious patients may suggest this.
Suggested management for dementia patients with sundowning includes use of structured activities at the relevant times (enforcement of external Zeitgebers) and increased staffing or availability of family members. Sedative medications are probably best avoided.

References

Cross References
Delirium; Dementia

Sunset Sign
- see SETTING SUN SIGN

Suppression
- see EXTINCTION

Supranuclear Gaze Palsy
A supranuclear gaze palsy results from pathology located above the cranial nerve nuclei supplying the extraocular muscles. Voluntary gaze is impaired while the integrity of the oculomotor nuclei and infranuclear connections may be demonstrated by the preservation of:

- Vestibulo-ocular reflexes (VOR): overcoming the ophthalmoplegia, at least in the early stages (e.g. the supranuclear gaze palsy in the vertical plane in progressive supranuclear palsy);
- Oculocephalic reflex (doll’s head, doll’s eye manoeuvre);
- Bell’s phenomenon.

Supranuclear gaze palsies may be classified as follows:

- **Horizontal:**
  - Hemisphere (frontal) lesion: eyes deviated to the side of the lesion, or in the case of an irritative (e.g. epileptic) focus away from the side of the lesion.
  - Paramedian pontine reticular formation: eyes deviated to contralateral side.

- **Vertical:**
  - Brainstem compression/distortion;
  - Dorsal upper midbrain (e.g. rostral interstitial nucleus of the median longitudinal fasciculus; pineal lesion causing Parinaud’s syndrome);
  - Skew deviation.

Recognized causes of supranuclear gaze palsy include

- Progressive supranuclear palsy (PSP; Steele–Richardson–Olszewski syndrome);
- Creutzfeldt–Jakob disease;
Swinging Flashlight Sign

- Corticobasal degeneration;
- Progressive subcortical gliosis of Neumann;
- Adult-onset Niemann–Pick disease;
- Gaucher’s disease.

Reference

Cross References
Gaze palsy; Parinaud’s syndrome; Parkinsonism; Prevost’s sign; Skew deviation; Vestibulo-ocular reflexes

Surface Dyslexia
- see ALEXIA

Suspended Sensory Loss
Sensory loss or impairment involving the trunk and proximal limbs may be described as suspended, or in a ‘cape-like’, ‘bathing suit’, ‘vest-like’, or cuirasse distribution. This may reflect intrinsic or intramedullary spinal cord pathology, in which case other signs of myelopathy may be present, including dissociated sensory loss, but it can also occur in peripheral neuropathic disease such as acute porphyria.

Cross References
Dissociated sensory loss; Myelopathy

Swan Neck
This term has been applied to thinning of the neck musculature, as in myotonic dystrophy type 1, for example.

Swearing
Swearing is not, in sensu strictu, a part of language, serving merely to add force of emotion to the expression of ideas; hence it is within the same category as loudness of tone or violence of gesticulation.

Reference

Cross Reference
Coprolalia

Sweat Level
A definable sweat level, below which sweating is absent, is an autonomic change which may be observed below a spinal compression.

Swinging Flashlight Sign
The swinging flashlight sign or test, originally described by Levitan in 1959, compares the direct and consensual pupillary light reflexes in one eye; the speed of swing is found by trial and error. Normally the responses are equal but in the
presence of an afferent conduction defect an inequality is manifest as pupillary dilatation. The test is known to be unreliable in the presence of bilateral afferent defects of light conduction. Subjective appreciation of light intensity, or light brightness comparison, is a subjective version of this test.

Reference

Cross References
Marcus Gunn pupil, Marcus Gunn sign; Pupillary reflexes; Relative afferent pupillary defect (RAPD)

Syllogomania
Syllogomania is excessive hoarding behaviour, often contributing to domestic squalor, as seen in Diogenes syndrome.

Synaesthesia
Synaesthesia is a perceptual experience in one sensory modality following stimulation of another sensory modality. The most commonly encountered example is colour-word synaesthesia (‘coloured hearing’ or chromaesthesia), experiencing a visual colour sensation on hearing a particular word. Synaesthesia occurs in a small percentage of the normal population. Known synaesthetes include the composers Messiaen and Scriabin, the artist Kandinsky, and the author Nabokov. There may be concurrent excellent memory (hypermnesia), sometimes of a photographic nature (eidetic memory). Symptomatic synaesthesia is rare but has been described with epileptic seizures of temporal lobe origin and with drug use (LSD). Characteristics ascribed to synaesthetic experience include its involuntary or automatic nature, consistency, generic or categorical and affect-laden quality.

Neuropsychologically, this phenomenon has been conceptualized as a break down of modularity. Functional imaging studies of colour-word synaesthetes show activation of visual associative areas of cortex (but not primary visual cortex), as well as perisylvian language areas, when listening to words which evoke the experience of colour.

References

Cross References
Auditory-visual synaesthesia; Phosphene

Synkinesia, Synkinesis
The term synkinesis may be used in different ways. It may refer to involuntary movements which accompany or are associated with certain voluntary
movements (*mitbewegungen*, motor overflow). These may be physiological, for example, the swinging of the arms when walking. Alternatively, such associated phenomena may be pathological, e.g. the involuntary contraction of orbicularis oculi when opening the mouth (the Marin–Amat syndrome: inverse Marcus Gunn phenomenon), acquired after lower motor neurone facial (VII) nerve palsies and presumed to reflect aberrant reinnervation. Aberrant nerve regeneration is common to a number of synkinetic phenomena, such as elevation of a ptotic eyelid on swallowing (Ewart phenomenon) and upper eyelid elevation or retraction on attempted downgaze (pseudo-Von Graefe’s sign). Crocodile tears, or lacrimation when salivating, due to reinnervation following a lower motor neurone facial nerve palsy, may also fall under this rubric, although there is no movement per se (autonomic synkinesis), likewise gustatory sweating.

Abnormal synkinesis may be useful in assessing whether weakness is organic or functional (cf. Hoover’s sign).

Synkinesis may also refer to the aggravation of limb rigidity detected when performing movements in the opposite limb (e.g. clenching and relaxing the fist), also known as activated rigidity or Froment’s sign.

**Cross References**

Crocodile tears; Ewart phenomenon; Froment’s sign; Gustatory sweating; Hoover’s sign; Jaw winking; Pseudo-Von Graefe’s sign; Rigidity
The ‘table top’ sign describes the inability to place the hand flat on a level surface, recognized causes of which include ulnar neuropathy (*main en griffe*), Dupuytren’s contracture, diabetic cheiroarthropathy, and camptodactyly.

**Cross Reference**

‘Prayer sign’

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**Tachylalia**

Tachylalia is increased speech velocity. This has been reported in patients with cerebrotendinous xanthomatosis, particularly in the 20–40-year age group.

**Reference**


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**Tachyphemia**

Tachyphemia is repetition of a word or phrase with increasing rapidity and decreasing volume; it may be encountered as a feature of the speech disorders in parkinsonian syndromes.

**Cross Reference**

Parkinsonism

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**Tactile Agnosia**

Tactile agnosia is a selective impairment of object recognition by touch despite (relatively) preserved somaesthetic perception. This is a unilateral disorder resulting from lesions of the contralateral inferior parietal cortex. Braille alexia may be a form of tactile agnosia, either associative or apperceptive.

**References**


**Cross Reference**

Agnosia

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**Tadpole Pupils**

Pupillary dilatation restricted to one segment may cause peaked elongation of the pupil, a shape likened to a tadpole’s pupil. This has been recorded in Horner’s syndrome, migraine, and Holmes–Adie pupil.
Talantropia

Reference

Talantropia
- see NYSTAGMUS

Tandem Walking
Tandem walking, or heel–toe walking, also known as the dynamic Romberg’s test, is the ability to walk along a straight line placing one foot directly in front of the other, heel to toe, which may be likened to walking a tightrope.

In ataxic disorders, cerebellar (midline cerebellum, in which axial coordination is most affected) or sensory (loss of proprioception), the ability to tandem walk is impaired, as reflected by the tendency of such patients to compensate for their incoordination by developing a broad-based gait.

Cross References
Ataxia; Cerebellar syndromes; Proprioception; Rombergism, Romberg’s sign

Tasikinesia
Tasikinesia is forced walking as a consequence of an inner feeling of restlessness or jitteriness as encountered in akathisia.

Reference

Cross Reference
Akathisia

Tay’s Sign
- see CHERRY RED SPOT AT THE MACULA

Teichopsis
Meaning literally ‘town-wall vision’, this term was coined by Airy in 1870 to describe the ‘bastioned form of transient hemiopsia’ which he experienced as part of his own migraine attacks, and illustrated in his paper.

Reference

Cross Reference
Fortification spectra

Telegraphic Speech
- see AGRAMMATISM

Telopsia
Telopsia is a visual illusion in which the image is altered in position; the term may also be used to refer to the image appearing abnormally distant (cf. porropsia).
Cross References
Illusion; Metamorphopsia; Pelopsia; Porropsia

Temporal Desaturation
Temporal desaturation refers to an impairment in perception of red targets confined to the temporal visual hemifield. This may be the earliest indication of a developing temporal field defect, as in a bitemporal hemianopia due to a chiasmal lesion, or a monocular temporal field defect (junctional scotoma of Traquair) due to a distal ipsilateral optic nerve lesion.

Cross References
Hemianopia; Scotoma

Temporal Pallor
Pallor of the temporal portion of the optic nerve head may follow atrophy of the macular fibre bundle in the retina, since the macular fibres for central vision enter the temporal nerve head. This may be associated with impairment of central vision.

Cross Reference
Optic atrophy

Terson Syndrome
Terson’s syndrome refers to vitreous haemorrhage in association with any form of intracranial or subarachnoid haemorrhage.

Tetanus, Tetany
- see MAIN D’ACCOUCHEUR; RISUS SARDONICUS; SPASM

Tetraparesis, Tetraplegia
- see QUADRIPARESIS, QUADRIPLEGIA

Threat Reflex
- see BLINK REFLEX

Tic
A tic is an abrupt, jerky repetitive movement involving discrete muscle groups, hence a less complex movement than a stereotypy. Vocal (phonic) tics are also described. Tics vary in intensity, lack rhythmicity, and are relatively easy to imitate. They may temporarily be voluntarily suppressed by will power (perhaps accounting for their previous designation as ‘habit spasms’) but this is usually accompanied by a growing inner tension or restlessness, only relieved by the performance of the movement.

The pathophysiology of tics is uncertain. The belief that Tourette syndrome was a disorder of the basal ganglia has now been superseded by evidence of dysfunction within the cingulate and orbitofrontal cortex, perhaps related to excessive endorphin release. The aetiological differential diagnosis of tic includes

- Idiopathic;
- Tourette syndrome;
- Tics related to structural brain damage;
• Drug-induced tics;
• Tics triggered by streptococcal infection.

Treatment of tics is most usually with dopamine antagonists (haloperidol, sulpiride) and opioid antagonists (naltrexone); clonidine (central α2 adrenergic-receptor antagonist) and tetrabenazine (dopamine-depleting agent) have also been reported to be beneficial on occasion. Botulinum toxin injections and deep brain stimulation have also been tried.

The word tic has also been used to describe the paroxysmal, lancinating pains of trigeminal neuralgia (tic douloureux).

References

Cross References
Klazomania; Stereotypy

Tic Convulsif

Tic convulsif is a name that has been given to the combination of trigeminal neuralgia (tic douloureux) with hemifacial spasm. Both may be characterized as neurovascular compression syndromes.

References

Tie Sign
- see VISUAL DISORIENTATION

Tilted Disc
Tilted optic disc is a benign anomaly, causing the discs (it is usually bilateral) to look oval or lopsided, with a bitemporal superior visual field defect which does not respect the vertical midline.

Reference

Cross References
Bitemporal hemianopia; Visual field defects

Tinel’s Sign (Hoffmann–Tinel Sign)
Tinel’s sign (Hoffmann–Tinel sign) is present when tingling (paraesthesia) is experienced when tapping lightly with a finger or a tendon hammer over a compressed or regenerating peripheral nerve. The tingling (Tinel’s ‘sign of formication’) is present in the cutaneous distribution of the damaged nerve (‘peripheral reference’). Although originally described in the context of peripheral nerve regeneration after injury, Tinel’s sign may also be helpful in diagnosing focal
entrapment neuropathy such as carpal tunnel syndrome. However, it is a ‘soft’ sign; like other provocative tests for carpal tunnel syndrome (e.g. Phalen’s sign) it is not as reliable for diagnostic purposes as electromyography (EMG). Its specificity has been reported to range between 23 and 60% and sensitivity between 64 and 87%.

A ‘motor Tinel sign’ has been described, consisting of motor EMG activity and jerking of muscles evoked by manipulation of an entrapped nerve trunk.

The neurophysiological basis of Tinel’s sign is presumed to be the lower threshold of regenerating or injured (demyelinated) nerves to mechanical stimuli, which permits ectopic generation of orthodromic action potentials, as in Lhermitte's sign.

References
D’Arcy CA, McGee S. Does this patient have carpal tunnel syndrome? JAMA 2000; 283: 3110–3117.

Cross References
Closed fist sign; Flick sign; Hand elevation test; Lhermitte’s sign; Phalen’s sign; Pressure provocation test

Tinnitus
Tinnitus is the perception of elementary non-environmental sound or noise in the ear. This is most usually a subjective phenomenon (i.e. heard only by the sufferer), occurring in the absence of acoustic stimulation. It may occur in conjunction with either conductive or sensorineural hearing loss. However, in about one-fifth of sufferers, tinnitus is objective (i.e. heard also by an observer). This may result from:

- **vascular causes**: e.g. arteriovenous malformation, fistula; carotid or vertebral bruit;
- **mechanical causes**: e.g. palatal myoclonus (ear click).

The common causes of subjective tinnitus are as follows:

- **middle/inner ear disease**: cochlear hydrops (Ménière’s disease), presbycusis, acoustic tumour;
- **pulsatile**: normal heartbeat, glomus jugulare tumour, raised intracranial pressure, cervical/intracranial aneurysm, arteriovenous malformation.

Cross References
Hallucination; Palatal myoclonus

‘Tip-of-the-Tongue’ Phenomenon
- see CIRCUMLOCUTION

Titubation
- see HEAD TREMOR
Todd’s Paralysis, Todd’s Paresis

Todd’s paralysis (Todd’s paresis) is a transient localized weakness (usually hemiparesis), lasting seconds to minutes (exceptionally 24–48 h), observed following a focal motor epileptic seizure or Jacksonian seizure originating in the central motor strip, or febrile convulsion, a phenomenon first described by RB Todd in 1854. The pattern and duration of postictal signs are quite heterogeneous. Aphasia is also described. A postictal ‘paralytic’ conjugate ocular deviation may be observed after adverse seizures. Todd’s paresis is of localizing value, being contralateral to the epileptogenic hemisphere.

The differential diagnosis of transient postictal hemiparesis includes stroke, hemiplegic migraine, and, in children, alternating hemiplegia.

References

Cross References
Hemiparesis; Seizures

Toe Walking

Toe walking, or cock walking, is walking on the balls of the toes, with the heel off the floor. A tendency to walk on the toes may be a feature of hereditary spastic paraplegia and the presenting feature of idiopathic torsion dystonia in childhood.

Cross Reference
Dystonia

Tongue Biting

Tongue biting is one feature of a seizure: in a generalized tonic–clonic epileptic seizure the side or sides of the tongue are typically bitten: a specific but not very sensitive sign. In a non-epileptic seizure the tip of the tongue may be more likely to be bitten.

Reference

Cross Reference
Seizure

Tonic Spasms

Painful tonic spasms occur in multiple sclerosis, especially with lesions of the posterior limb of the internal capsule or cerebral peduncle, perhaps due to ephaptic activation, or following putaminal infarction.

References
Transcortical Aphasias


**Cross Reference**
Spasm

**Torpor**
- see OBTUNDATION

**Torticollis**
Torticollis (wryneck, cervical dystonia, nuchal dystonia, spasmodic torticollis) is a movement disorder characterized by involuntary contraction of neck musculature, involving especially sternocleidomastoid, trapezius, and splenius capitis. In the majority of cases (>50%) this produces head rotation, but laterocollis, retrocollis, tremulous (‘no–no’) and complex (i.e. variable) forms are seen; antecollis is unusual. Contractions are usually unilateral, may be associated with local pain, and, as with other types of dystonia, may be relieved by a ‘sensory trick’ (*geste antagoniste*). Causes of torticollis include

- Idiopathic (the majority);
- Secondary to acquired cervical spine abnormalities, trauma;
- Cervical spinal tumour;
- Tardive effect of neuroleptics.

The treatment of choice is botulinum toxin injections into the affected muscles. Injections benefit up to 70–80% of patients, but need to be repeated every 3 months or so.

**Reference**

**Cross References**
Antecollis; Dystonia; *Geste antagoniste*; Laterocollis; Retrocollis

**Tortopia**
- see ENVIRONMENTAL TILT

**Tourettism**
- see TIC

**Transcortical Aphasias**
Transcortical aphasias may be categorized as either motor or sensory.

- **Transcortical motor aphasia (TCMA):**
  There is a dissociation between preserved repetition (cf. conduction aphasia) and impaired fluency, manifest as delayed initiation, even mutism, impaired lexical selection, and reduced capacity to generate unconstrained syntactic forms. TCMA is associated with pathology (usually infarction) in the supplementary motor area, superior to Broca’s area (left lateral frontal cortex) or in subcortical structures including white matter projections and dorsal caudate nucleus; it has clinical similarities with Broca’s aphasia.
Transcortical sensory aphasia (TCSA):
There is a dissociation between preserved repetition (cf. conduction aphasia) and impairments of spoken and written language comprehension without phonemic paraphasia. TCSA is associated with pathology (usually infarction) in the ventral and ventrolateral temporal lobe involving the fusiform gyrus and the inferior temporal gyrus, and posterior convexity lesions involving the posterior middle temporal gyrus and the temporooccipital junction. It has similarities with Wernicke’s aphasia.

Some authorities prefer to label these conditions as ‘extrasylvian aphasis syndromes’, to distinguish them from the perisylvian aphasis syndromes (Broca, Wernicke, conduction); moreover, these syndromes are not ‘transcortical’ in any literal sense.

Dynamic aphasia may be a lesser version of TCMA, in which there are no paraphasias and minimal anomia, preserved repetition, and automatic speech, but reduced spontaneous speech. This may be associated with lesions of dorsolateral prefrontal cortex (‘frontal aphasis’) in the context of frontal lobe degeneration. There may be incorporational echolalia, when the patient uses the examiner’s question to help form an answer.

References

Cross References
Aphasia; Broca’s aphasis; Conduction aphasis; Dynamic aphasis; Echolalia; Paraphasia; Wernicke’s aphasis

Transverse Smile
- see ‘MYASTHENIC SNARL’

Tremblement Affirmatif, Tremblement Negatif
- see HEAD TREMOR

Tremor
Tremor is an involuntary movement, roughly rhythmic and sinusoidal, although some tremors (e.g. dystonic) are irregular in amplitude and periodicity. Tremors may be classified clinically:

- Rest tremor:
  present when a limb is supported against gravity and there is no voluntary muscle activation, e.g. the 3.5–7 Hz ‘pill rolling’ hand tremor of Parkinson’s disease; midbrain/rubral tremor.

- Action tremor:
  present during any voluntary muscle contraction.
Various subtypes of action tremor are recognized:

- **Postural tremor:** present during voluntary maintenance of a posture opposed by gravity, e.g. arm tremor of essential tremor; 6 Hz postural tremor sometimes seen in Parkinson’s disease (‘re-emergent tremor’), which may predate emergence of akinesia/rigidity/rest tremor; modest postural tremor of cerebellar disease; some drug-induced tremors (including alcohol withdrawal, delirium tremens); tremor of IgM paraproteinaemic neuropathy; wing-beating tremor of Wilson’s disease.

- **Kinetic tremor:** present with movement, often with an exacerbation at the end of a goal-directed movement (intention tremor), e.g. cerebellar/midbrain tremor (3–5 Hz).

- **Task-specific tremor:** evident only during the performance of a highly skilled activity, e.g. primary writing tremor.

- **Isometric tremor:** present when voluntary muscle contraction is opposed by a stationary object, e.g. primary orthostatic tremor (14–18 Hz).

- **Psychogenic tremors:** these are difficult to classify, with changing characteristics; the frequency with which such tremors are observed varies greatly between different clinics; the coactivation sign (increase in tremor amplitude with peripheral loading) is said to be typical of psychogenic tremor.

EMG may be useful for determining tremor frequency, but is only diagnostic in primary orthostatic tremor.

Various treatments are available for tremor, with variable efficacy. Essential tremor often responds to alcohol, and this is a reasonable treatment (previous anxieties that such a recommendation would lead to alcoholism seem unjustified); alternatives include propranolol, topiramate, primidone, alprazolam, flunarizine, and nicardipine. In Parkinson’s disease, tremor is less reliably responsive to levodopa preparations than akinesia and rigidity; anticholinergics such as benzhexol may be more helpful (but may cause confusion). Primary orthostatic tremor has been reported to respond to gabapentin, clonazepam, primidone, and levodopa. Cerebellar tremor is often treated with isoniazid, but seldom with marked benefit, likewise carbamazepine, clonazepam, ondansetron, limb weights; stereotactic surgery may be an option in some patients disabled with tremor.

**References**


**Cross References**
Asterixis; Coactivation sign; Head tremor; Holmes’ tremor; Knee tremor; Palatal tremor; Parkinsonism; Vocal tremor, Voice tremor; Wing-beating tremor

**Trendelenburg’s Sign**
Trendelenburg’s sign is tilting of the pelvis towards the side of the unaffected raised leg in a unilateral superior gluteal nerve lesion.

**Triparesis**
- see SEQUENTIAL PARESIS

**Triplopia**
Triplopia is seeing triple, a rare complaint, which may be due to oculomotor nerve palsy, internuclear ophthalmoplegia, and abducens nerve palsy. It may result from an unusual interpretation of abnormal eye movements

**Reference**

**Cross References**
Abducens (VI) nerve palsy; Diplopia; Internuclear ophthalmoplegia (INO); Oculomotor (III) nerve palsy

**Trismus**
Trismus is an inability to open the jaw due to tonic spasm or contracture of the masticatory muscles, principally masseter and temporalis, effecting forced jaw closure (‘lockjaw’). Recognized causes and associations of trismus include
- Dystonia of the jaw muscles (e.g. drug-induced dystonic reaction);
- Generalized tonic–clonic epileptic seizure;
- Neuromuscular diseases: polymyositis, tetanus, nemaline myopathy, trauma to the muscles of mastication, rabies, strychnine poisoning;
- Infection in the pterygomandibular space;
- Metabolic disorders: Gaucher’s disease (type II);
- Central disorders: brainstem encephalopathy, multiple sclerosis, pseudobulbar palsy.

**Reference**

**Cross References**
Dystonia; Pseudobulbar palsy

**Trombone Tongue**
Trombone tongue, or flycatcher tongue, refers to an irregular involuntary darting of the tongue in and out of the mouth when the patient is requested to keep the tongue protruded. This sign may be seen in choreiform movement disorders such as Huntington’s disease and neuroacanthocytosis and in tardive dyskinesia.
Cross References
Chorea, Choreoathetosis; Impersistence; Milkmaid’s grip

Trömner’s Sign
Trömner’s sign is flexion of the thumb and index finger in response to tapping or flicking the volar surface of the distal phalanx of the middle finger, held partially flexed between the examiner’s finger and thumb. This is an alternative method to Hoffmann’s sign (‘snapping’ the distal phalanx) to elicit the finger flexor response. As in the latter, it is suggestive of a corticospinal tract (upper motor neurone) lesion above C5 or C6, especially if unilateral, although it may be observed in some normal individuals.

Cross References
Hoffmann’s sign; Upper motor neurone (UMN) syndrome

Trousseau’s Sign
Trousseau described the signs and symptoms of tetany, including anaesthesia, paraesthesia, and the main d’accoucheur posture, as well as noting that the latter could be reproduced by applying a bandage or inflating a cuff around the arm so as to impede circulation; the latter is now known as Trousseau’s sign and indicates latent tetany.

Trousseau also noted the concurrence of venous thrombosis and migrating thrombophlebitis with malignant disease, also referred to as Trousseau’s sign; this may present with cerebral venous thrombosis.

Reference

Cross References
Achromatopsia; Chvostek’s sign; Main d’accoucheur

Tullio Phenomenon
The Tullio phenomenon is the experience of vestibular symptoms and signs (vertigo, nystagmus, oscillopsia, postural imbalance, ocular tilt reaction, +/− skew deviation) on exposure to high-intensity acoustic stimuli, presumed to be due to hyperexcitability of the normal vestibular response to sound, causing pathological stimulation of the semicircular canals and/or otoliths. This unusual phenomenon may be associated with perilymph leaks or a defect in the capsule forming the roof of the anterior semicircular canal. The sound sensitivity is probably at the level of the receptors rather than the vestibular nerve.

Reference

Cross References
Nystagmus; Ocular tilt reaction; Oscillopsia; Skew deviation; Vertigo

Tunnel Vision
A complaint of ‘tunnel vision’ may indicate constriction of the visual field. This may be observed with enlargement of the blind spot and papilloedema as a
consequence of raised intracranial pressure or with a compressive optic neuropathy. The normal visual field enlarges the further away from the eye the visual target used to map the field is held, hence there is in fact ‘funnel vision’. In non-organic visual impairment, by contrast, the visual field stays the same size with more distant targets (tunnel vision).

A tunnel vision phenomenon has also been described as part of the aura of seizures of anteromedial temporal and occipitotemporal origin. A closing in of vision may be described as a feature of presyncope.

**Cross References**
Aura; Blind spot; Hemianopia; Papilloedema; Visual field defects

**Two-Point Discrimination**
Two-point discrimination is the ability to discriminate two adjacent point stimuli (e.g. using a pair of callipers) as two rather than one. The minimum detectable distance between the points (acuity) is smaller on the skin of the fingertips (i.e. greater acuity) than, say, the skin on the back of the trunk. Impairments of two-point discrimination may occur with dorsal column spinal cord lesions, in which proprioception (and possibly vibration) is also impaired. Cortical parietal lobe lesions may produce a cortical sensory syndrome of astereognosis, graphaesthesia, and impaired two-point discrimination.

**Cross References**
Astereognosis; Graphaesthesia; Proprioception; Vibration
Uhthoff’s Phenomenon
Uhthoff’s phenomenon or symptom is the worsening of visual acuity (‘amblyopia’ in Uhthoff’s 1890 description) with exercise in optic neuritis, reflecting the temperature sensitivity of demyelinated axons (i.e. reduced safety factor for faithful transmission of action potentials). The term has subsequently been applied to exercise and/or temperature related symptoms in other demyelinated pathways. It has also been described in the context of other optic nerve diseases, including Leber’s hereditary optic neuropathy, sarcoidosis, and tumour.

Evidence suggesting that Uhthoff’s phenomenon is associated with an increased incidence of recurrent optic neuritis, and may be a prognostic indicator for the development of multiple sclerosis, has been presented.

Inverse Uhthoff sign, improved vision with warming, has been described.

References

Cross References
Lhermitte’s sign; Phosphenes

Unterberger’s Sign
Unterberger’s sign or Unterberger’s stepping test is said to examine the integrity of vestibulospinal connections and attempts to define the side of a vestibular lesion. The patient is asked to march on the spot with the eyes closed (i.e. proprioceptive and visual cues are removed); the patient will rotate to the side of a unilateral vestibular lesion (Unterberger’s sign). The test is not very useful, particularly in chronic, progressive, or partially compensated vestibular lesions.

Reference

Cross References
Proprioception; Vertigo
Upper Motor Neurone (UMN) Syndrome
An upper motor neurone (UMN) syndrome constitutes a constellation of motor
signs resulting from damage to upper motor neurone pathways, i.e. proximal
to the anterior horn cell. These may be termed ‘pyramidal signs’, but since
there are several descending motor pathways (e.g. corticospinal, reticulospinal,
vestibulospinal), of which the pyramidal or corticospinal pathway is just one,
‘upper motor neurone syndrome’ is preferable. ‘Long tract signs’ may be a more
accurate term, often used interchangeably with ‘pyramidal signs’. The syndrome
may be variable in its clinical features but common elements, following the
standard order of neurological examination of the motor system, include the
following:

- **Appearance:**
  Usually normal, but there may be wasting in chronic UMN syndromes,
  but this is usually not as evident as in lower motor neurone syndromes;
  contractures may be evident in chronically spastic limbs.

- **Tone:**
  Hypertonus, with spasticity, clasp-knife phenomenon and sustained
  clonus.

- **Power:**
  Weakness, often in a so-called pyramidal distribution (i.e. affecting
  extensors more than flexors in the upper limb, and flexors more
  than extensors in the lower limb); despite its clinical utility, the term
  pyramidal is, however, a misnomer (see Weakness).

- **Coordination:**
  Depending on the degree of weakness, it may not be possible to
  comment on the integrity of coordination in UMN syndromes; in a
  pure UMN syndrome coordination will be normal, but syndromes
  with both ataxia and UMN features do occur (e.g. spinocerebellar
  syndromes, ataxic hemiparesis syndromes).

- **Reflexes:**
  Limb hyperreflexia, sometimes with additional reflexes indicative of
corticospinal tract involvement (Hoffmann’s sign, Trömner’s sign,
crossed adductor reflex); Babinski’s sign (extensor plantar response);
and cutaneous reflexes (abdominal, cremasteric) are lost.

  The most reliable (‘hardest’) signs of UMN syndrome are increased tone,
  clonus, and upgoing plantar responses. The most subtle (i.e. earliest) sign of
  UMN involvement is debated but may be pronator drift or impaired forearm
  and finger rolling.

  The clinical phenomena comprising the upper motor neurone syndrome
  may be classified as ‘positive’ and ‘negative’ depending on whether they reflect
  increased or decreased activity in neural pathways:
Urinary Retention

- **Positive:**
  - Exaggerated stretch/tendon reflexes, flexor spasms;
  - Clonus;
  - Autonomic hyperreflexia;
  - Contractures.

- **Negative:**
  - Muscle weakness, pronator drift;
  - Loss of dexterity.

These features help to differentiate UMN from LMN syndromes, although clinically the distinction is not always easy to make: a ‘pyramidal’ pattern of weakness may occur in LMN syndromes (e.g. Guillain–Barré syndrome) and acute UMN syndromes may cause flaccidity and areflexia (e.g. ‘spinal shock’).

**Reference**

**Cross References**
Abdominal reflexes; Ataxic hemiparesis; Babinski’s sign (1); Clasp-knife phenomenon; Clonus; Contracture; Cremasteric reflex; Forearm and finger rolling; Hoffmann’s sign; Hyperreflexia; Hypertonia, Hypertonus; Lower motor neurone (LMN) syndrome; Pronator drift; Pseudobulbar palsy; Spasticity; Trömner’s sign; Weakness

**Urgency**
- see INCONTINENCE

**Urinary Retention**
Although urinary retention is often urological in origin (e.g. prostatic hypertrophy) or a side effect of drugs (e.g. anticholinergics), it may have neurological causes. It may be a sign of acute spinal cord compression, with or without other signs in the lower limbs, or of acute cauda equina compression, for example, with a central L1 disc herniation. Sometimes the level of the pathology is several segments above that expected on the basis of the (‘false localizing’) neurological signs. Loss of awareness of bladder fullness may lead to retention of urine with overflow.

A syndrome of urinary retention in young women has been described, associated with myotonic-like activity on sphincter EMG; this condition may be associated with polycystic ovary disease and is best treated with clean intermittent self-catheterization.

**References**

**Cross References**
Cauda equina syndrome; ‘False localizing signs’; Incontinence; Myelopathy; Paraplegia; Radiculopathy
Useless Hand of Oppenheim

The deafferented hand or arm is functionally useless and manifests involuntary movements due to severe proprioceptive loss. This was first described in multiple sclerosis by Oppenheim in 1911 and reflects plaques in the dorsal root entry zone of the relevant spinal cord segment(s).

References

Cross References
Proprioception; Pseudoathetosis; Pseudochoreoathetosis

Utilization Behaviour
Utilization behaviour is a disturbed response to external stimuli, a component of the environmental dependency syndrome, in which seeing an object implies that it should be used. Two forms of utilization behaviour are described:

- **Induced:**
  When an item is given to the patient or their attention is directed to it, e.g. handing them a pair of spectacles which they put on, followed by a second pair, which are put on over the first pair.

- **Incidental or spontaneous:**
  When the patient uses an object in their environment without their attention being specifically directed towards it.

  Another element of the environmental dependency syndrome which coexists with utilization behaviour is imitation behaviour (e.g. echolalia, echopraxia). Primitive reflexes and hypermetamorphosis may also be observed.

  Utilization behaviour is associated with lesions of the frontal lobe, affecting the inferior medial area bilaterally. It has also been reported following paramedian thalamic infarction.

References

Cross References
Automatic writing behaviour; Echolalia; Echopraxia; Frontal lobe syndromes; Hypermetamorphosis; Imitation behaviour; Primitive reflexes
Valsalva Manoeuvre
The Valsalva manoeuvre is a simple test of autonomically mediated cardiovascular reflexes, comprising forced expiration against resistance (‘straining’), followed by release of the resistance and completion of expiration. The first phase produces impaired cardiac filling due to impaired venous return as a consequence of elevated intrathoracic pressure, with a fall in cardiac output and blood pressure, inducing peripheral vasoconstriction (sympathetic pathways) to maintain blood pressure. The second phase causes a transient overshoot in blood pressure as the restored cardiac output is ejected into a constricted circulation, followed by reflex slowing of heart rate.

In autonomic (sympathetic) dysfunction, reflex vasoconstriction, blood pressure overshoot, and bradycardia do not occur. The latter may be conveniently assessed by measuring R–R intervals in a prolonged ECG recording, an R–R interval ratio between the training and release phases of less than 1.1 suggesting impaired baroreceptor response.

Cross Reference
Orthostatic hypotension

Vegetative States
The vegetative state is a clinical syndrome in which cognitive function is lost, due to neocortical damage (hence no awareness, response, speech), whilst vegetative (autonomic, respiratory) function is preserved due to intact brainstem centres. Primitive postural and reflex limb movements may also be observed. The syndrome, also known as neocortical death, coma vigil, and the apallic syndrome, may be seen after extensive ischaemic–hypoxic brain injury, for example, following resuscitation after cardiac arrest, and needs to be distinguished from coma, akinetic mutism, and the locked-in syndrome. Persistent vegetative state (PVS) is defined by persistence of this state for > 12 months (UK) or > 6 months (USA) after brain trauma, or > 6 months (UK) or > 3 months (USA) following brain anoxia. The prognosis of PVS is poor, but occasional reports of very late recovery have appeared.

References

Cross References
Akinetic mutism; Coma; Locked-in syndrome
Vertigo is an illusion of movement, a sense of rotation or of tilt, causing a feeling of imbalance or disequilibrium. It is a subtype of ‘dizziness’, to be distinguished from the light-headedness of general medical conditions (vasovagal attacks, presyncope, cardiac dysrhythmias). Vertigo is often triggered by head movement and there may be associated autonomic features (sweating, pallor, nausea, vomiting). Vertigo may be horizontal, vertical, or rotatory.

Pathophysiologically, vertigo reflects an asymmetry of signalling anywhere in the central or peripheral vestibular pathways. Clinically, it may be possible to draw a distinction between central and peripheral lesions: in the latter there may be concurrent hearing loss and tinnitus (reflecting vestibulocochlear (VIII) nerve involvement). Facial weakness (VII) and ipsilateral ataxia suggest a cerebellopontine angle lesion; diplopia, bulbar dysfunction, and long tract signs are suggestive of a central pathology. Peripheral vertigo tends to compensate rapidly and completely with disappearance of nystagmus after a few days, whereas central lesions compensate slowly and nystagmus persists.

The clinical pattern of vertigo may give clues as to underlying diagnosis:

<table>
<thead>
<tr>
<th>Vertigo</th>
<th>Peripheral</th>
<th>Central</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acute</td>
<td>Labyrinthitis</td>
<td>Brainstem/cerebellum haemorrhage/infarct/demyelination</td>
</tr>
<tr>
<td>Prolonged, spontaneous</td>
<td>Otomastoiditis, Vestibular neur(on)itis</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Labyrinthine concussion</td>
<td></td>
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<tr>
<td></td>
<td>Isolated labyrinthine infarct</td>
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<tr>
<td></td>
<td>Vestibular nerve section</td>
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<td></td>
<td>Drug-induced</td>
<td></td>
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<tr>
<td>Recurrent, episodic</td>
<td>Migraine</td>
<td>Vertebrobasilar ischaemia (with associated features)</td>
</tr>
<tr>
<td></td>
<td>Menière’s disease (endolympathic hydrops)</td>
<td></td>
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<tr>
<td></td>
<td>Autoimmune inner ear disease (isolated, systemic)</td>
<td></td>
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<tr>
<td></td>
<td>Perilymph fistula</td>
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<tr>
<td></td>
<td>Epilepsy (rare)</td>
<td></td>
</tr>
<tr>
<td>Positional</td>
<td>Benign paroxysmal positional vertigo (BPPV)</td>
<td>Fourth ventricle lesions: multiple sclerosis, Chiari malformation, brainstem/cerebellar tumours</td>
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<tr>
<td></td>
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<td>Spino cerebellar atrophy</td>
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<td></td>
<td></td>
<td>Neurological disorder</td>
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<td></td>
<td>Psychogenic</td>
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<tr>
<td>Chronic</td>
<td>Vestibular decompensation/failure</td>
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</tbody>
</table>
All patients with vertigo should have a Hallpike manoeuvre performed during the examination.

Specific treatments are available for certain of these conditions. A brief course of a vestibular sedative (cinnarizine, Serc) is appropriate in the acute phase, but exercises to ‘rehabilitate’ the semicircular canals should be begun as soon as possible in peripheral causes. In BPPV, most patients respond to the Epley manoeuvre to reposition the otoconia which are thought to cause the condition (canalolithiasis). Brandt–Daroff exercises are an alternative. Cawthorne–Cooksey exercises are helpful in vestibular decompensation or failure.

References

Cross References
Ataxia; Caloric testing; Facial paresis; Hallpike manoeuvre, Hallpike test; Hennebert’s sign; Illusion; Nystagmus; Vestibulo-ocular reflexes

Vestibulo-ocular Reflexes
The vestibulo-ocular reflexes (VORs) are physiological mechanisms to generate eye rotations that compensate for head movements, especially during locomotion, so stabilizing the retinal image on the fovea. VORs depend upon the integrity of the connections between the semicircular canals of the vestibular system (afferent limb of reflex arc) and oculomotor nuclei in the brainstem (efferent limb). Loss of vestibular function, as in acute bilateral vestibular failure, causes gaze instability due to loss of VORs, causing the symptom of oscillopsia when the head moves. As well as vestibular input, compensatory eye rotations may also be generated in response to visual information (pursuit–optokinetic eye movements) and neck proprioceptive information; anticipatory eye movements may also help stabilize the retinal image.

VORs are also useful in assessing whether ophthalmoplegia results from a supranuclear or infranuclear disorder, since in the former the restriction of eye movement may be overcome, at least in the early stages, by the intact VOR, e.g. the supranuclear gaze palsy in the vertical plane in progressive supranuclear palsy.

VORs are difficult to assess in conscious patients because of concurrent pursuit–optokinetic eye movements and because rotation of the head through large angles in conscious patients leads to interruption of VORs by vestibular nystagmus in the opposite direction (optokinetic nystagmus). The head impulse test may be used to test VORs in conscious patients, for example, those with vertigo in whom vestibular failure is suspected. VOR may also be assessed using a slow (0.5–1.0 Hz) doll’s head manoeuvre whilst directly observing the eyes (‘catch up’ saccades may be seen in the absence of VOR), by measuring visual
acuity (dynamic visual acuity, or illegible E test; dropping two to three lines on visual acuity with head movement vs. normal if VOR impaired), and by ophthalmoscopy (optic disc moves with head if VOR abnormal).

In unconscious patients, slow phase of the VORs may be tested by rotating the head and looking for contraversive conjugate eye movements (oculocephalic responses, doll’s head eye movements) or by caloric testing. VORs are lost in brainstem death.

Another important element of VOR assessment is suppression or cancellation of VOR by the pursuit system during combined head and eye tracking. VOR suppression may be tested by asking the patient to fixate on their thumbs with arms held outstretched whilst rotating at the trunk or sitting in a swivel chair. VOR suppression can also be assessed during caloric testing: when the nystagmus ceases with fixation, removal of the fixation point (e.g. with Frenzel’s glasses) will lead to recurrence of nystagmus in normals but not in those with reduced or absent VOR suppression. VOR suppression is impaired (presence of nystagmus even with slow head movements) in cerebellar and brainstem disease.

References

Cross References
Caloric testing; Coma; Doll’s eye manoeuvre, Doll’s head manoeuvre; Hallpike manoeuvre, Hallpike test; Head impulse test; Ocular tilt reaction; Oculocephalic response; Oscillopsia; Supranuclear gaze palsy; Vertigo

Vibration
Vibratory sensibility (pallaesthesia) represents a temporal modulation of tactile sense. On this ground, some would argue that the elevation of vibration to a ‘sensory modality’ is not justified. Vibratory sensibility is easily tested using a tuning fork (128 Hz). This assesses the integrity of rapidly adapting mechanoreceptors (Pacinian corpuscles) and their peripheral and central connections; the former consist of large afferent fibres, the latter consist of ascending projections in both the dorsal and lateral columns. The classification of both vibration and proprioception as ‘posterior column signs’, sharing spinal cord and brainstem pathways, is common in neurological parlance (and textbooks) but questioned by some. Instances of dissociation of vibratory sensibility and proprioception are well recognized, for instance the former is usually more impaired with intramedullary myelopathies.

Decrease in sensitivity of vibratory perception (increased perceptual threshold) is the most prominent age-related finding on sensory examination, thought to reflect distal degeneration of sensory axons.

References

**Cross References**
Age-related signs; Myelopathy; Proprioception; Two-point discrimination

**Visual Agnosia**
Visual agnosia is a disorder of visual object recognition. The term derives from Freud (1891), but it was Lissauer (1890), speaking of *seelenblindheit* (psychic blindness), who suggested the categorization into two types:

- **Apperceptive visual agnosia:**
  A defect of higher-order visual perception leading to impaired shape recognition, manifested as difficulty copying shapes or matching shapes, despite preserved primary visual capacities, including visual acuity and fields (adequate to achieve recognition), brightness discrimination, colour vision, and motion perception (indeed motion may facilitate shape perception; see Riddoch’s phenomenon). Reading is performed with great difficulty, with a ‘slavish’ tracing of letters which is easily derailed by any irrelevant lines; such patients may appear blind.

- **Associative visual agnosia:**
  An impairment of visual object recognition thought not to be due to a perceptual deficit, since copying shapes of unrecognized objects is good. The scope of this impairment may vary, some patients being limited to a failure to recognize faces (prosopagnosia) or visually presented words (pure alexia, pure word blindness).

These terms continue to be used, although some authors (e.g. Critchley) have taken the view that there is always some qualitative or quantitative disorder of sight, and hence that to isolate subtypes is a ‘vain pursuit’.

Visually agnosic patients can recognize objects presented to other sensory modalities. Clinically, apperceptive visual agnosia lies between cortical blindness and associative visual agnosia.

Apperceptive visual agnosia results from diffuse posterior brain damage; associative visual agnosia has been reported with lesions in a variety of locations, usually ventral temporal and occipital regions, usually bilateral but occasionally unilateral. Pathological causes include cerebrovascular disease, tumour, degenerative dementia (visual agnosia may on occasion be the presenting feature of Alzheimer’s disease, the so-called visual variant, or posterior cortical atrophy), and carbon monoxide poisoning. A related syndrome which has on occasion been labelled as apperceptive visual agnosia is simultanagnosia, particularly the dorsal variant in which there is inability to recognize more than one object at a time. Associative visual agnosia has sometimes been confused with optic aphasia.

**References**

Visual Disorientation

Visual disorientation refers to the inability to perceive more than a fragment of the visual field at any one time; it is sometimes characterized as a shifting fragment or island of clear vision. There may be difficulty fixating static visual stimuli and impaired visual pursuit eye movements.

Visual disorientation may be demonstrated by sitting directly opposite to the patient and asking them, whilst looking at the bridge of the examiner’s nose, to reach for the examiner’s hand held up in the peripheral field of vision. Once contact is made with the hand, the examiner holds up the other hand in a different part of the field of vision. Individuals with visual disorientation will find it hard to see the hand and will grope for it, sometimes mistakenly grasping the examiner’s clothing (‘tie sign’) or face.

Visual disorientation is secondary to, and an inevitable consequence of, the attentional disorder of dorsal simultanagnosia, in which the inability to attend two separate loci leads to impaired localization. It may be a feature of Alzheimer’s disease; indeed, sometimes it may be the presenting feature, but there are usually signs of more generalized cognitive problems (e.g. impairment of episodic memory). Visual disorientation may be localizing to the non-dominant hemisphere.

References

Cross References
Simultanagnosia; Visual agnosia

Visual Extinction

Visual extinction is the failure to respond to a novel or meaningful visual stimulus on one side when a homologous stimulus is given simultaneously to the contralateral side (i.e. double simultaneous stimulation), despite the ability to perceive each stimulus when presented singly.

Cross References
Extinction; Neglect

Visual Field Defects

Visual fields may be mapped clinically by confrontation testing. The most sensitive method is to use a small (5 mm) red pin, more so than a waggling finger. Peripheral fields are tested by moving the target in from the periphery, and the patient asked to indicate when the colour red becomes detectable, not when they
first see the pinhead. The central field may be mapped using the same target presented statically to points within the central field.

The exact pattern of visual field loss may have localizing value due to the retinotopic arrangement of fibres in the visual pathways: any unilateral area of restricted loss implies a prechiasomatic lesion (choroid, retina, optic nerve), although lesions of the anterior calcarine cortex can produce a contralateral monocular temporal crescent. Bilateral homonymous scotomata are postchiasmal in origin; bilateral heteronymous scotomata may be seen with chiasmal lesions. Topographically, typical visual field defects are as follows:

- **Retina**: monocular visual loss, altitudinal field defects; central or centrocaecal scotoma, ring scotoma;
- **Optic nerve**: central or centrocaecal scotoma; junctional scotoma of Traquair;
- **Optic chiasm**: bitemporal hemianopia; junctional scotoma;
- **Optic tract**: homonymous hemianopia, usually incongruous;
- **Lateral geniculate nucleus**: homonymous hemianopia, usually incongruous;
- **Optic radiations**: homonymous hemianopia, usually congruous; quadrantanopia;
- **Visual cortex**: homonymous hemianopia, usually congruous; quadrantanopia; cortical blindness.

**References**

**Cross References**
Altitudinal field defect; Hemianopia; Junctional scotoma, Junctional scotoma of Traquair; Macula sparing, Macula splitting; Quadrantanopia; Scotoma; Tilted disc

**Visual Form Agnosia**
This name has been given to an unusual and a highly selective visual perceptual deficit, characterized by loss of the ability to identify shape and form, although colour and surface detail can still be appreciated, but with striking preservation of visuomotor control (i.e. a pattern of deficits inverse to those seen in optic ataxia). This syndrome is thought to reflect selective damage to the ventral (‘what’) stream of visual processing in the lateral occipital area, whilst the dorsal (‘where’) stream remains intact, yet the workings of the latter are not available to consciousness.

**Reference**

**Cross References**
Agnosia; Optic ataxia; Visual agnosia
Visual Grasp Reflex
- see SACCADES

Visual Perseveration
Visual perseveration may refer to more than one unusual subjective visual experience:

- Hallucinatory and recurring appearance of an object after its removal: palinopsia (q.v.);
- Visual perseveration in sensu strictu, when a disappearing visual stimulus does not fade from view; no recurrence as in palinopsia;
- Visual stimulus sensed over an unduly extensive area of environmental space: visuospatial perseveration or illusory visual spread; rare; no temporal factor, when the stimulus is removed the effect disappears.

References

Cross References
Palinopsia; Perseveration

Visuopalpebral Reflex
- see BLINK REFLEX

Vocal Tremor, Voice Tremor
Vocal or voice tremor is a shaking, quivering, or quavering of the voice. It may be heard in:

- Essential tremor;
- Cerebellar disorders;
- Spasmodic dysphonia/laryngeal dystonia;
- Parkinson’s disease;
- Motor neurone disease.

The pathophysiology is uncertain but may relate to rhythmic contractions of the cricothyroid and rectus abdominis muscles.

Cross References
Dysphonia; Tremor

Von Graefe’s Sign
Von Graefe’s sign, or Graefe’s sign, is the retarded descent of the upper eyelid during movement of the eye from the primary position to downgaze; the lid ‘follows’ the eye. This may be termed ‘lid lag’, although some authorities reserve this term for a static situation in which the lid is higher than the globe on downgaze. Von Graefe’s sign may be seen in thyroid ophthalmopathy.

Cross References
Lid lag; Pseudo-Von Graefe’s sign
Vulpian's Sign

Vorbereiden
- see GANSER PHENOMENON

VOR Suppression
- see VESTIBULO-OCULAR REFLEXES

Vulpian's Sign
- see PREVOST'S SIGN
Waddling Gait
Weakness of the proximal leg and hip girdle muscles, most often of myopathic origin, impairs the stability of the pelvis on the trunk during walking, leading to exaggerated rotation with each step, an appearance likened to the waddling of a duck. In addition, the hips may be slightly flexed and lumbar lordosis exaggerated. Neurogenic causes include spinal muscular atrophy and Guillain–Barré syndrome.
Cross Reference
Myopathy

‘Waiter’s Tip’ Posture
Lesions of the upper trunk of the brachial plexus (Erb–Duchenne type) produce weakness and sensory loss in the C5 and C6 distribution, typically with the arm hanging at the side, internally rotated at the shoulder with the elbow extended and the forearm pronated: the ‘waiter’s tip’ posture, also sometimes known as the ‘porter’s tip’ or ‘policeman’s tip’.
Cross References
Plexopathy; Radiculopathy

Wallenberg’s Syndrome
- see LATERAL MEDULLARY SYNDROME

Wall-Eyed
- see EXOTROPIA; INTERNUCLEAR OPHTHALMOPLEgia (INO)

Warm-Up Phenomenon
Easing of muscle stiffness with repeated contraction, the warm-up phenomenon, is reported by many patients with myotonia congenita (Thomsen’s disease, Becker’s disease), in contrast to the situation in paramyotonia.
Cross References
Myotonia; Paramyotonia

Wartenberg’s Sign (1)
In ulnar neuropathy, Wartenberg’s sign refers to the slightly greater abduction of the fifth digit on the affected side, due to paralysis of the adducting palmar interosseous muscle and unopposed action of the radial-innervated extensor muscles (digitii minimi, digitorum communis).
Cross Reference
Froment’s sign
Wartenberg’s Swing Test
Wartenberg’s swing test is used to assess limb and trunk rigidity (cf. pendulum test of Wartenberg, used to measure spasticity, q.v.). With the patient standing, the examiner holds the shoulders and gently shakes backwards and forwards, the two sides out of phase. Normally, the passive arm swing induced by this movement will be out of phase with the trunk movements, but in rigidity the limbs and trunk tend to move en bloc. Passive swinging of the wrist or elbow joint may also be performed to assess rigidity.

Cross References
Parkinsonism; Rigidity; ‘Rolex’ sign; Spasticity

Wasting
Wasting refers to a thinning of the musculature, also known as atrophy or, if of neurogenic origin, amyotrophy. Wasting may be a consequence of disorders of:
- muscle (myopathies, dystrophies);
- peripheral nerve (more so in axonal than demyelinating peripheral neuropathies);
- anterior horn cells (e.g. motor neurone disease).

Wasting may occur in chronic upper motor neurone syndromes (e.g. chronic hemiplegia) but is not as evident as in lower motor neurone syndromes where wasting may appear subacutely (over a few weeks).

Wasting may also be seen in general medical disorders associated with a profound catabolic state, e.g. cancer cachexia, uncontrolled heart failure, liver cirrhosis, and renal failure.

Cross References
Amyotrophy; Atrophy; Lower motor neurone (LMN) syndrome; Upper motor neurone (UMN) syndrome

Weakness
Weakness is an objective loss of muscle strength. This is conveniently quantified or rated using the MRC grading system:
- 5 = normal power;
- 4 = active movement against gravity and resistance;
- 3 = active movement against gravity;
- 2 = active movement with gravity eliminated;
- 1 = flicker or trace of contraction;
- 0 = no contraction (paralysis).

However, this is not a linear scale; grade 4 often becomes subdivided into 4−, 4, and 4+ (or even 5−) according to the increasing degree of resistance which the examiner must apply to overcome activity. It is also important to assess what effort the patient is making to comply with the testing; ‘apparent weakness’ or ‘pseudoparesis’ may be shorthand for lack of patient effort. Sudden ‘giving way’ of muscle contraction may be an indicator of this. Non-uniform resistance may also be due to pain (algesic pseudoparesis). Testing records only
the best-forced maximal contraction and should not develop into an unseemly trial of strength between patient and examiner. Accepting all these difficulties, it should be acknowledged that the grading of weakness, like all clinical observations, is subject to some degree of observer bias.

The pattern of muscle weakness may suggest its anatomical origin. So-called pyramidal weakness (i.e. affecting upper limb extensors more than flexors and lower limb flexors more than extensors) suggests an upper motor neurone lesion (corticospinal pathways). However, there is no evidence that pure lesions of the pyramidal tracts produce this picture: pyramidotomy in the monkey results in a deficit in fine finger movements, but without weakness. Moreover, a similar pattern of weakness may be observed in lower motor neurone disorders such as Guillain–Barré syndrome. Coexistent wasting suggests that muscle weakness is of lower motor neurone origin, especially if acute, although wasting may occur in long-standing upper motor neurone lesions. Weakness with minimal or no muscle wasting may be non-organic, but may be seen in conditions such as multifocal motor neuropathy with conduction block.

Reference
Aids to the examination of the peripheral nervous system. London: HMSO, 1976.

Cross References
Collapsing weakness; Hyperreflexia; Lower motor neurone (LMN) syndrome; Upper motor neurone (UMN) syndrome; Wasting

Weber’s Test
Weber’s test is one of the tuning fork tests, which may be used to confirm a conductive component in unilateral or asymmetric hearing loss. The vibrating tuning fork (512 Hz preferred) is put on the middle of the forehead and the patient asked in which ear it is heard; this depends entirely upon bone conduction (BC). Hence the sound localizes to the side of a conductive hearing loss (where bone conduction is greater than air conduction, BC > AC) and away from the side of a sensorineural hearing loss.

Reference

Cross Reference
Rinne’s test

Wernicke’s Aphasia
Wernicke’s aphasia is the classical ‘receptive aphasia’, in distinction to the ‘expressive aphasia’ of Broca, although this classification is problematic since there are concurrent ‘expressive’ problems in Wernicke’s aphasia. Other terms sometimes used for Wernicke-type aphasia are sensory aphasia or posterior aphasia.

Considering each of the features suggested for the clinical classification of aphasias (see Aphasia), Wernicke’s aphasia is characterized by:

- **Fluency**: fluent speech with phonemic and semantic paraphasias and para-grammatism (inappropriate use of syntax); ‘empty speech’ with few verbs and nouns; prosody usually preserved; at worst, flowing speech (logorrhoea)
devoid of semantic meaning (jargon aphasia, semantic aphasia); automatic speech is often better preserved than spontaneous, e.g. counting, days of week, overlearned phrases (‘I’m fine’);

- **Comprehension**: impaired auditory comprehension (sine qua non; ‘word deafness’); impaired reading comprehension probably also required (not specifically discussed by Wernicke);
- **Repetition**: impaired;
- **Naming**: severely impaired (anomia) and not aided by cueing (cf. Broca’s aphasia);
- **Reading**: usually impaired, with numerous paralexic errors and impaired reading comprehension (cf. pure word deafness);
- **Writing**: similarly affected.

There may be associated anxiety, with or without agitation and paranoia, and concurrent auditory agnosia. Because of a loss of self-monitoring of output, patients are often not aware of the impairment, and behavioural disturbance sometimes misdiagnosed as ‘acute confusional state’ and even referral or admission to psychiatric hospital may occur, particularly if there is no or minimal accompanying hemiparesis. The differential diagnosis of Wernicke’s aphasia includes delirium and schizophrenia.

The neuroanatomical substrate of Wernicke’s aphasia has been a subject of debate. Wernicke placed it in the posterior two-thirds of the superior temporal gyrus and planum temporale (Brodmann area 22), but more recent neuroradiological studies (structural and functional imaging) suggest that this area may be more associated with the generation of paraphasia, whereas more ventral areas of temporal lobe and angular gyrus (Brodmann areas 37, 39, and 40) may be associated with disturbance of comprehension. A correlation exists between the size of the lesion and the extent of the aphasia. A similar clinical picture may occur with infarcts of the head of the left caudate nucleus and left thalamic nuclei.

**References**

**Cross References**
Agnosia; Agraphia; Alexia; Anomia; Aphasia; Broca’s aphasia; Jargon aphasia; Logorrhoea; Paraphasia; Pure word deafness; Schizophrenia; Transcortical aphasias

**Wheelchair Sign**
The so-called wheelchair sign describes patients with parkinsonism who start to use a wheelchair for mobility early in the course of their disease, usually because of repeated falls. Early falls are a typical feature of progressive supranuclear palsy (Steele–Richardson–Olszewski syndrome), but not idiopathic Parkinson’s disease or other parkinsonian syndromes.

**Cross References**
Parkinsonism; ‘Rocket sign’
Woltman's Sign

Wing-Beating Tremor
Wing-beating tremor is absent at rest but develops when the arms are extended, hence this is a postural tremor. It is said to be typical of Wilson's disease (hepatolenticular degeneration).

Cross Reference
Tremor

Winging of the Scapula
Winging of the scapula, or scapula alata, is a failure to hold the medial border of the scapula against the rib cage when pushing forward with the hands. It is most easily observed by asking the patient to push or press against a wall or the examiner’s hand whilst observing the scapula which lifts away from the posterior chest wall.

Winging of the scapula may be a consequence of weakness of the serratus anterior muscle, usually due to a neuropathy of the long thoracic nerve of Bell, but sometimes as a consequence of brachial plexus injury or cervical root (C7) injury. It may also be of myopathic origin, as in facioscapulohumeral dystrophy.

Weakness of trapezius, particularly the middle trapezius muscle, may also cause winging of the upper part of the scapula, more prominent on abduction of the arm, when the superior angle of the scapula moves farther from the midline. Hence spinal accessory (XI) nerve palsy enters the differential diagnosis.

Witzelsucht
Witzelsucht, or the joking malady, refers to excessive and inappropriate facetiousness or jocularity, a term coined in the 1890s for one of the personality changes observed following frontal (especially orbitofrontal) lobe injury. This phenomenon may overlap with those described as moria or emotional lability.

Cross References
Emotionalism, Emotional lability; Frontal lobe syndromes; Moria

Woltman’s Sign
Woltman’s sign denotes slow-relaxing, or ‘hung-up’, tendon reflexes. These are most commonly seen in the context of untreated hypothyroidism, but have also been recorded in other situations, including treatment with β-blockers, diabetes mellitus, and complete heart block. The phenomenon is sometimes labelled ‘pseudomyotonia’ because of its superficial resemblance to the slow muscle relaxation of myotonia, but electrophysiological testing does not show myotonic discharges.

Chorea may result in apparently ‘hung-up’ reflexes, perhaps due to a choreiform jerk after muscle relaxation.

The mechanisms underlying Woltman’s sign are uncertain: changes in basal metabolic rate and in muscle fibre types (selective loss of fast twitch fibres) have been suggested.

References
Wrestler’s Sign

This name has been given to the excessive effort in irrelevant muscle groups accompanied by prominent non-verbal signs of effort such as grunting in patients with apparent (‘functional’) weakness. It may coexist with intermittent voluntary effort, collapsing weakness, cocontraction of agonist and antagonist muscles, and inconsistency in clinical examination (e.g. inability to lift leg from couch when recumbent, despite preserved ability to stand up and walk).

Cross Reference
Collapsing weakness

Wrist Drop

Wrist drop describes a hand hanging in flexion due to weakness of wrist extension. This results from radial nerve palsy, either in the axilla or spiral groove of the humerus (‘Saturday night palsy’, although other nerves may also be compressed by hanging the arm over a chair, e.g. ulnar, median). Distal lesions affecting branches of the posterior interosseous branch of the radial nerve may produce more circumscribed deformity, such as weak extension of metacarpophalangeal joints (‘finger drop’, ‘thumb drop’).

Writer’s Cramp

Writer’s cramp, also known as graphospasm, la crampe des écrivains, or scrivener’s palsy, is a focal dystonia involving the hand and/or arm muscles, causing abnormal posturing of the hand when writing; it is the most common of the task-specific dystonias (once known as ‘craft palsies’). When attempting to write, patients may find they are involuntarily gripping the pen harder, and there may also be involuntary movement at the wrist or in the arm. A tremor may also develop, not to be confused with primary writing tremor in which there is no dystonia. Handwriting becomes illegible. Attempts to use the contralateral hand may be made, but this too may become affected with time (‘mirror dystonia’). The problem may be exclusive to writing (simple writer’s cramp) but some people develop difficulties with other activities as well (e.g. shaving; dystonic writer’s cramp), reflecting a dystonia of the hand or arm. Muscle fatigue may make writing more legible. Familial forms may be associated with mutations in the epsilon sarcoglycan gene (DYT11). There may be an association between writer’s cramp and carpal tunnel syndrome.

There is some neurophysiological evidence that the condition is due to abnormalities within the spinal cord segmental motor programmes and muscle spindle afferent input to them.

Writer’s cramp may be amenable to local botulinum toxin injections into the hand or arm muscles responsible for the involuntary movement. Other strategies which may be used include writing with a different grip (e.g. whole hand grip), using a fat-bodied pen or using a word processor.
References

Cross References
Dystonia; Fatigue; Tremor

Wrong-Way Eyes
- see PREVOST’S SIGN

Wry Neck
- see TORTICOLLIS
Yawning

Yawning is an arousal reflex thought to be generated in the brainstem reticular formation to counteract brain hypoxia; it may precede vasovagal syncope. Excessive or pathological yawning (chasm) is compulsive, repetitive yawning not triggered by physiological stimuli such as fatigue or boredom. Known associations of excessive yawning or salvos of yawning include

- Presyncope
- Hypoglycaemia
- Drugs: SSRIs, imipramine, valproate, dopamine agonists
- Migraine prodrome
- Temporal lobe epileptic seizures
- Encephalitis
- Multiple sclerosis
- Tumours of the fourth ventricle, frontal lobes
- Electroconvulsive therapy
- Postthalamotomy
- Neuroleptic withdrawal
- Parkinson’s disease, progressive supranuclear palsy, restless legs syndrome
- Pseudobulbar palsy of motor neurone disease

Although the mechanisms are uncertain, yawning may represent a disturbance of dopaminergic neurotransmission. Levodopa may help.

References

Cross References
Parkinsonism; Sighing

Yips

Yips is the name given to a task-specific focal dystonia seen in golfers, especially associated with putting.

Reference

Cross Reference
Dystonia
Yo-Yo-ing
Yo-yo-ing is a form of dyskinesia experienced by patients with idiopathic Parkinson’s disease who have been treated for several years with levodopa preparations, in which there are sudden and unpredictable swings between hypokinesia/akinesia (‘off’ state; freezing) and severe hyperkinesia (‘on’ state), sometimes known as the ‘on–off phenomenon’. Yo-yo-ing is difficult to treat: approaches include dose fractionation, improved drug absorption, or use of dopaminergic agonists with concurrent reduction in levodopa dosage.

Reference

Cross References
Akinesia; Dyskinesia; Hypokinesia
Zeitraffer Phenomenon
The zeitraffer phenomenon has sometimes been described as part of the aura of migraine, in which the speed of moving objects appears to increase, even the vehicle in which the patient is driving.

Reference

Zooagnosia
The term zooagnosia has been used to describe a difficulty in recognizing animal faces. This may be observed as a component of prosopagnosia. In one case, this deficit seemed to persist despite improvement in human face recognition, suggesting the possibility of separate systems for animal and human face recognition; however, the evidence is not compelling. In a patient with developmental prosopagnosia seen by the author, there was no subjective awareness that animals such as dogs might have faces.

References

Cross References
Agnosia; Prosopagnosia

Zoom Effect
The zoom effect is a metamorphopsia occurring as a migraine aura in which images increase and decrease in size sequentially.

Cross Reference
Metamorphopsia