NEUROLOGICAL INVESTIGATIONS

Imaging of the spinal cord

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Methods of investigation

HISTORICAL PERSPECTIVE

Air myelography

The first contrast medium used to show the spinal cord was air. The first reports of its use to localise intraspinal tumours came from Jacobeus in 1921 and Dandy in 1925.1 The technique was later refined to show the entire spinal canal; this involved complete replacement of the CSF by air and distension of the spinal subarachnoid space.2 Adequate visualisation of the spinal cord usually required tomography. Spinal roots were not shown, and many types of pathology such as vascular malformations or arachnoiditis were either not shown or easily misinterpreted. Despite the fact that the technique was difficult to achieve and very hard on the patient, it remained the preferred method for visualising the spinal cord in many centres until the advent of non-toxic water soluble contrast only just over 10 years ago.

Oil myelography

At the time air myelography was being developed, it was found accidentally that iodised oils could be moved through the spinal subarachnoid space under the influence of gravity. They proved easier to use than air, and this quickly established oil myelography as the technique of choice especially in the lumbar spinal canal. It received even greater impetus from the appearances in 1940, of iophendylate (Myodil; Pantopaque), a preparation that was less viscous than the earlier Lipiodol and better for demonstrating the spinal cord. Myodil was very opaque to x rays and special techniques such as tomography were not required. It was very slowly absorbed and could be left in the canal and rerun postoperatively to check the adequacy of decompression. Its main disadvantage was immiscibility with CSF; it tended to break up into globules, forming a layer in the spinal canal, which made it difficult to demonstrate both anterior and posterior surfaces of the spinal cord unless large amounts were used. For nearly 40 years, Myodil was generally the agent of choice, with air being reserved for special situations, such as spinal dysraphism and syringomyelia. As late as 1989, eminent names in neurosurgical surgery were still declaring their preference for Myodil, mainly because of experiences with non-diagnostic water soluble myelograms in which the contrast medium had become too dilute. Myodil is now no longer manufactured, and existing stocks have been withdrawn because of the frequency with which it caused chronic adhesive arachnoiditis.3,4

WATER SOLUBLE MYEOGRAPHY

The advantages of water soluble over oily contrast media were established by use of a substance called Abrodil in Scandinavian countries. This provided superior images of the cauda equina and root sheaths. Other ionic water soluble media such as Conray and Dimer-X enjoyed limited use, but all were too toxic to use other than to show the lower lumbar thecal sac. They are, therefore, irrelevant to the present review. A revolution in myelography occurred when the new non-ionic water soluble medium metrizamide (Amipaque) appeared. This was far less neurotoxic than previous ones. It could be used safely around the spinal cord. Inadvertent deposition in the head when running the contrast medium into the cervical region often caused generalised seizures, however; a risk minimised by introducing it by lateral C1–2 puncture.5

By 1983, similar and even less toxic media had been developed and metrizamide was quickly withdrawn. These agents, such as iohexol, do not cause arachnoiditis in the concentrations used in clinical practice, and by 1989, Skalpe and Sortland were able to state that “epileptic seizures have not been reported following myelography with Omnipaque [iohexol], so it seems that fear of this complication can be virtually disregarded.”6

These media are still not perfect: most patients experience postmyelography headache and about 5% become confused or develop symptoms such as radicular pain or meningism.7 Further agents have been developed, but clinical trials are now difficult to mount because myelography is so little used. Iohexol should be injected by lumbar puncture wherever possible to reduce the risk of injury to the spinal cord or vertebral artery in lateral cervical puncture.8 Myelography may cause neurological deterioration due to the spinal puncture, causing injury to the spinal cord or intraspinal bleeding; the disease, resulting in increased cord compression during
the positioning required for radiography; the procedure, causing raised intraspinal pressure below a subarachnoid block.1,8

**X Ray Computed Tomography (CT) of the Spine**

Soon after its introduction for cranial imaging CT was used in the spine. It provided the valuable cross sectional perspective of the spinal canal, formerly very difficult to achieve with conventional radiography. Artefacts caused by bone degrade intraspinal contrast in spinal CT. Intravenous contrast media increase contrast between extradural tissues and CSF, and intrathecal contrast media provide excellent visualisation of the spinal cord and other intradural structures. Patients requiring myelography usually are booked for CT as well, the findings on the myelogram or the clinical features serving to direct the CT examinations to specific levels. Computed tomography is almost exclusively a cross sectional technique; sagittal and coronal projections require reformatting from stacks of axial slices, and resolution in reformatted planes is not as good as in the plane of data acquisition. Radiation dose can be very considerable, and new guidelines for the use of CT in Britain have been published recently.9

**Magnetic Resonance Imaging (MRI)**

During the period when myelography was being greatly improved, MRI appeared, and within just a few years myelography was pushed almost into obsolescence. With modern high resolution MRI, almost all intradural features demonstrable by myelography can be shown, usually better, by MRI. Myelography is now indicated only when satisfactory MRI cannot be obtained because it is contraindicated (pacemakers, mechanical heart valves, aneurysm clips); it cannot be done (claustrophobia and anaesthesia refused or unsafe, patient cannot fit into the magnet due to obesity, scoliosis, or limb contractions); or it is not available quickly enough, due to lack of on call service or other logistical problems.

Several recent developments have had a special impact on MRI of the spinal cord.

**Volumetric (3D) Acquisitions**

The deployment of fast image techniques has permitted three dimensional spatial encoding within a few minutes. This results in multiple contiguous images, no interslice gaps, and section thicknesses down to 1 or 2 mm. The best images of the spinal cord structure are, however, still usually obtained from thicker slices. The best results from volumetric MRI acquisitions are usually obtained on high-field machines.

**Fast Spin Echo (FSE)**

Several phase encoding steps are made at each excitation instead of just one, which greatly reduces acquisition times and permits the use of much larger matrices. This results in twice the resolution in even shorter data acquisition periods. The penalty is slight loss of contrast and increased sensitivity to physical motion, for which it is more difficult to compensate than when single phase encoding steps are used.

**Phased Array Coils**

Spinal imaging requires surface coils, and the phased array configuration enables data to be acquired simultaneously from two or more surface coils. This permits imaging of the entire spinal cord over one acquisition period and greatly reduces imaging time. Vertebral level counting, sometimes difficult or impossible from single coils, especially in the thoracic region, also becomes easy.

A wide variety of postprocessing options are available on most imagers, or can be purchased separately. These permit multplanar reformatting in real time, three dimensional surface rendering, colour coding, and many other modifications. The ability to reconstruct in a curved plane is potentially useful in scoliosis, but we have found this of only limited value because most major curvatures are in more than one plane.

** Artefacts**

Artefacts are important to consider because they can closely simulate intraspinal disease. Phase dispersion across the image, due to magnetic susceptibility variation and chemical shift effects, reduces sensitivity to biological signal differences and reduces boundary definition. The truncation artefact, generated at boundaries by image processing, is particularly relevant at the CSF-spinal cord interface, and is one possible cause of the band of high or low signal seen in the centre of the cord in midsagittal images. It also causes difficulty in defining the position of the cord-CSF boundary where the problem is compounded by susceptibility effects. Phantom studies have shown that both electronic and caliper measurements of the spinal cord, especially in the phase encoding direction, can be artificially reduced by over 2 mm and may create a spurious impression of spinal cord flattening.10

**Motion Artefacts**

Motion artefacts are generated by cardiosynchronous and oscillatory motion of CSF. This motion has been documented and roughly quantified by MRI. At C2/3, movement is estimated at about 0·65 ml per cardiac cycle, downwards on systole and upwards on diastole.11 New data confirm older work indicating that the primary driving force behind intracranial and spinal canal CSF flow is expansion of the brain during vascular systole. The spinal cord and brain stem also descend very slightly on systole and oscillate anteroposteriorly with CSF flow.12 In classic studies, Rubin et al13 showed how the oscillatory motions generate linear artefacts parallel to the cord-CSF interface at roughly harmonic intervals across the images in the phase encoding direction, producing signal variation in the spinal cord image that could be easily misinterpreted as intramedullary pathology.13 Areas of turbulent CSF flow in...
regions where subarachnoid septa exist, particularly in the thoracic spine, can result in signal variations simulating intradural masses or enlarged vessels. Many strategies have been developed to minimise these problems, but results can be less consistent than is desirable.

The thoracic cord in particular is difficult to image well, especially in cross section, where cardiac motion and the proximity of the aorta add to the motion generated artefacts.

**Metallic artefacts**

Metallic artefacts can be particularly destructive of image quality. Many spinal operations utilise metal stabilisation devices, such as plates, rods, screws, and loops, and patients often require postoperative imaging at some stage. Ferromagnetic substances such as stainless steel generate major local artefacts and usually render spinal imaging useless. Tiny fragments from drills and punches, invisible on plain radiographs, may also result in devastating artefacts.\(^\text{14, 15}\) Design of implants also can be important such as the avoidance of conductive loops.\(^\text{14}\) The use of titanium for manufacture may have advantages.\(^\text{17}\) A wide range of titanium devices has become available only recently, however, and the long-term stability and biological effects of these new materials remain under evaluation.\(^\text{18}\)

**Special techniques**

New methods of brain imaging are usually eventually applied to the spine. Many are being evaluated and only some will be given passing reference here.

**Phase contrast imaging** utilises bipolar phase encoding gradients; the first brings all spins into phase, the second records reduced signal from moving spins as they dephase relative to stationary spins, due to their motion. This can be used to demonstrate molecular diffusion. One method displays apparent diffusion coefficients across the image and fluid filled cavities appear much brighter than solid areas.\(^\text{19}\) Phase contrast imaging can also be used to demonstrate coherent CSF flow and movement of the neural axis. **Phase contrast cine MR** produces images that represent velocity as a function of time throughout the cardiac cycle, and by the direction of the phase shift indicate flow direction.\(^\text{20}\) **Spatial modulation of magnetism (SPAMM)**, also referred to as presaturation bolus tracking, is a method whereby regions are tagged by applying a narrow band of saturation before the pulse sequence. This produces a dark stripe across the image that bends in the direction of movement. Multiple presaturation bands can be used to produce a “zebra stripe” pattern, and coupling with a cine loop increases sensitivity in detecting minimal movement.\(^\text{21}\) This has been used extensively to study syringomyelia. **Susceptibility contrast weighted dynamic MRI** has been used to study blood flow in spinal tumours and arteriovenous malformations.\(^\text{22, 23}\) A susceptibility weighted gradient recalled echo sequence is used to detect the large, transient signal reduction that occurs on the first pass of a bolus dose of intraventricular gadolinium through the lesion. **Fluid attenuated inversion recovery (FLAIR)** sequences are said to offer improved lesion detectability by permitting heavily T2 weighted acquisition to be accomplished with suppression of all signal from CSF. This removes motion artefact from CSF, which appears paradoxically as signal void on heavily T2 weighted images. Cysts and cystic lesions within the cord also appear, however, as signal void. Most other pathology appears as hyperintensity. At present, imaging with FLAIR is slow and of low resolution,\(^\text{24, 25}\) and like most of the functional imaging methods listed, it has found little general clinical application at present.

**SPINAL SONOGRAPHY**

**Intraoperative spinal sonography** has been used widely in some centres. A minimum of a two or three level laminectomy wound is required, filled with water or saline to act as an acoustic window.\(^\text{26, 27}\) Although considerable utility has been achieved,\(^\text{23}\) experience has shown that it may not distinguish tumour from cord tissue, may incorrectly identify cysts as solid masses due to unusual echogenicity of some cyst fluids, and cannot reliably distinguish tumoral from non-tumoral cysts.\(^\text{28}\) Surprisingly large transverse excursion of the spinal cord and roots is observed normally, lagging slightly behind the cardiac cycle, and breathing and the Valsalva manoeuvre produce additional abrupt movement. **Percutaneous spinal sonography** is possible in infants.\(^\text{27, 28}\) The spinal canal can also be imaged in adults by angling the probe parallel to an intervertebral disc space. **Transoesophageal transducers** have been used with some success to image the thoracic cord, but may be non-existent with the availability of MRI. **Transuterine sonography** has been used to identify dysraphic states in the fetus,\(^\text{29}\) but MRI may also be employed for this purpose.

**PLAIN RADIOGRAPHS OF THE SPINE**

Plain films have little part to play in the investigation of spinal cord disease. Inferences can be made about possible sites of spinal cord compression in conditions such as spondylotic myelopathy and trauma, and may suggest the site and type of cord involvement in dysraphic states. The role of plain films in the preliminary investigation of patients with intraspinal lesions was reviewed by Naidich et al as late as 1986.\(^\text{30}\) The spinal canal is usually enlarged in children with intramedullary tumours. Greatest sensitivity is achieved by graphing the interpediculate distances and comparing them with normal values, but even then the false positive rate is at least 11%. Therefore, plain radiographs have only a very limited screening value even in children.

**SPINAL ANGIOGRAPHY**

The technique of spinal angiography was developed over 30 years ago by Djindjian and
Doppman. Today it is indicated only for localisation of the major radiculomedullary arteries before operations on the spine, to study vascular malformations associated with arteriovenous shunts, and before endovascular treatment of a variety of spinal lesions. To demonstrate all the spinal cord arteries, selective injections into the intercostal, lumbar, lumbosacral, vertebral, deep, and ascending cervical arteries is required. Searches for dural fistulae may also require selective injections into branches of the external carotid artery. General anaesthesia is desirable although not essential. Moderate, usually transient, neurological deterioration occurs not uncommonly after extensive spinal angiography,11 but paraplegia is rare.12 Spinal angiography can be a very laborious undertaking; short cuts have been devised using other imaging modalities and these will be discussed with vascular malformations.

Shape, internal structure, and biomechanics of the spinal cord

SIZE AND SHAPE

Spondylotic flattening of the formalin fixed spinal cord is commonly found at necropsy, and was often considered to be a fixation artefact. Modern imaging has made it abundantly clear that this is not so. Good qualitative agreement is found between the appearance of the spinal cord on cross sectional imaging in living subjects and formalin fixed cords at necropsy,13 14 but clinical measurement is problematic. The most robust measure of size is cross sectional area. Area measurements could theoretically be accurate within about ± 5%,15 but on clinical images it is clear that nothing like this is actually achieved. Electronic window settings profoundly influence all measurements of the cord on computed myelography16 and on MRI many other factors are also involved. This is shown by the mean values derived by various workers for their control populations. To give three examples: mean normal cord cross sectional areas at C2 (where spondylosis is not expected) of 62 mm²,17 86-6 mm²,18 and 110 mm²19 have been reported. Close correlations between area measurements made by computed myelography and MRI were reported by Fukushima et al.10 (x = 0-901)41; but the mean values were very different—namely, 0-38 (SD 0-14) cm² on computed myelography and 0-50 (0-16) cm² on MRI. Yu and his colleagues concluded that each department needed to define its own normal range.42 It is hardly surprising that no workers have shown a stable relation between cord size, patient age, or body size. Few measurements have been published for the thoracic cord, but we can console ourselves that medical science is not too much the poorer for that. The two most commonly used methods of cord shape have been used: circularity (4π area/circumference)15 and the compression ratio (ratio of the anteroposterior to transverse diameters).15 40 41 Abnormal shapes have also been classified qualitatively in specific types of cord compression, especially cervical spondylisis and its variants, the most comprehensive being by Yu et al.10

INTERNAL STRUCTURE

The internal structure of the spinal cord on MRI seems remarkably similar to an anatomical preparation stained for myelin. More myelinated regions generally yield a lower signal than less myelinated ones. Magnetic resonance imaging has consistently shown variations in texture in histologically uniform regions, such as the anterior horns in the sacral area, Clarke’s column, and the dorsal horn complex; the gracile fascicles yield a slightly higher signal than the cuneate.42 43 In some types of image, the subpial zone has yielded a high signal44 and it is uncertain if this represents an MR artefact, or the narrow band of subpial degeneration commonly seen in the cords of aging subjects. It is notable that some authors have forgotten the T1 shortening that occurs with formalin fixation, making it difficult to obtain images with T1 weighted contrast.

BIOMECHANICAL PROPERTIES

Deformation of the cord by transverse compression of up to about 20% requires minimal force, whereas deformation in excess of 50% requires forces exceeding capillary perfusion pressure and begins to disrupt both transverse and longitudinal axons.45 46 Moreover, spinal cord substance has only a limited capacity for elastic recoil. It has been shown by measurements of cord deformation between flexion and extension on computed myelograms that only about 20% of a deformation recovers elastically.47 This has an important implication for interpreting CT and MRI images that are generally performed in positions where the available space for the spinal cord is maximised. Because of its lack of elasticity, it is safe to conclude that if the spinal cord is normal in cross sectional shape, it is not being appreciably compressed in any situation occurring in the patient during normal daily activities; and when deformity is present, it is safe to conclude that the deformity shown reasonably represents the magnitude of intermittent compression. We developed the simple concept of congruous cord deformity, to help distinguish deformation due to intermittent compression from that due to atrophy. When the available subarachnoid space appears capacious on cross sectional images, compression is suggested, nevertheless, if the deformity of the spinal cord is reciprocally congruous with the visible (disc/osteophyte) or expected (reducible subluxation) deformity of the spinal canal; in cord atrophy the deformation is incongruous.48

The compressed spinal cord usually increases in size after operative decompression. Two simple methods10 measured a mean postoperative increase in size of the compressed spinal cord of 13%, varying between 5% and 20% in different subgroups of patients, which is of the order one would expect from cord elasticity studies. The cord
also often changes shape after decompression, becoming less flattened or altered in some other way, because it is very easily deformed by varying conditions.

CORD MOTION
The cord lies about 2 mm more posteriorly in the supine than in the prone position, and the cord and vertebral midlines differ by up to 2 mm in over 40% of non-scoliotic subjects.49 On phase contrast MRI the range of normal cardiosynchronous oscillatory longitudinal movement of the upper cord has been measured to be only 0-4-0-5 mm.12 Initial studies caused excitement because they suggested that assessment of tethering could be made without objective influence of morphological appearances. Reduced oscillatory motion has been seen in the presence of cord tethering, and increased oscillatory motion in Chiari associated syringomyelia.21 The actual relevance of any of these findings, however, in the absence of morphological changes, such as a low lying conus medullaris, has, in my opinion, yet to be demonstrated.

Pathological states in cord substance
Necrosis
Water soluble contrast medium in the subarachnoid space diffuses freely into neural tissue, and x ray attenuation of spinal cord substance measured on CT equilibrates dynamically at about 20% of that in the subarachnoid space, being removed by the capillary-venous system.10 When necrosis occurs, hydrophobic lipid is broken down and the capillary bed destroyed. Contrast medium continues to accumulate in the necrotic region until the CSF concentration falls, and passive diffusion out of the area takes several hours. Necrotic areas thus appear as circumscribed areas of contrast enhancement on computed myelograms, often most conspicuous after six hours.50

On MRI, colliquative necrosis appears as a circumscribed area of signal change. Necrosis, from whatever cause, most often involves mainly the central parts of the spinal cord, in particular the grey matter and ventral parts of the posterior columns. Involvement is usually bilateral, producing either a localised confluent lesion or, more often, bilateral lesions resulting in an appearance likened to snakes' eyes on cross sectional images.52

WALLERIAN DEGENERATION
Antegrade degeneration in the long tracts of the spinal cord is seen with pathological states that cause axonal damage. Typically it appears as descending degeneration in the anterior part of the lateral column and ascending degeneration in the posterior columns, which, in diseases of the cervical spine, is often most severe in the fasciculus cuneatus.31 44 53

Wallerian degeneration is shown only by MRI. Its appearances have been studied mostly in the brain stem, and only recently in the spinal cord. Four stages are distinguishable on imaging, which evolve over about 14 weeks.59 For up to four weeks, MRI is normal and the chronic stage, characterised by volume loss and increased MR signal, persists indefinitely. Enhancement with intravenous contrast agents does not occur at any stage. In the spinal cord only the chronic stage has been described,44 55 but recent experimental studies with magnetisation transfer imaging have demonstrated abnormalities in the earliest stage.16

CYSTIC DEGENERATION AND SYRINGOMYELIA
Syringomyelia may result from any pathological process that is liable to cause spinal cord necrosis. It represents an end stage that itself may be progressive and promote further cord damage. The cavities are usually located dorsal to the central canal, with which they may or may not communicate. They may be single or multiple. Only about 15% extend beyond C2, and those that do so usually bifurcate around the decussations in the medulla oblongata and come to lie ventral to the floor of the fourth ventricle. Hydromyelia is a cavity consisting mainly of a dilated central canal and communicating with the fourth ventricle. It is closely associated with hydrocephalus, and is collapsed by ventricular shunting.57

Chiari I associated syringomyelia
In the currently most widely accepted hydro-mechanical theory of causation, this begins as a hydromyelia that subsequently loses communication with the fourth ventricle early in life, resulting in the condition often referred to as syringohydromyelia. Obstruction of the foramen magnum or the outlets of the fourth ventricle by the ascended cerebellar tonsils is central to most hydromechanical theories. In support, phase contrast cine MRI has demonstrated absence of CSF in the cisterna magna due to the abnormal cerebellar tonsils, and restoration of CSF flow after foramen magnum decompression, accompanied by collapse of the syrinx. Also, two groups have recently separately reported a new finding on dynamic MRI: accentuated caudal displacement of the cerebellar tonsils and spinal cord with cardiac systole, which was restored to the normal range, or obliterated altogether, by decompression of the foramen magnum.50 58 Oldfield et al proposed intermittent piston like obstruction of the foramen magnum by the tonsils.58

Although these mechanisms may operate in cases where the cerebellar tonsils do obstruct the foramen magnum, other workers have shown that the foramen magnum is not obstructed by the abnormal cerebellar tonsils in at least 20% of cases, and the cerebellum would need to move more than 10 times as far as has been recorded to cause intermittent obstruction.59 60 Furthermore, no association exists between the distension or cranial extent of the syrinx and the presence or absence of foramen magnum obstruction, or degree of tonsillar descent; indeed syringomyelia occurs significantly more often with mild rather than with severe tonsillar descent.60 61
Additional complications have arisen with respect to the nature of the Chiari I malformation itself. Firstly, assessment of the level of the cerebellar tonsils on midline sagittal MR images has greatly overestimated the prevalence of tonsillar descent below the foramen magnum. Indeed, a prevalence of between 15% and 20% is still generally accepted despite the previous assessments on myelography and computed cisternography indicating a prevalence of less than 1%. The recent volumetric MRI study of Savvy et al, however, has confirmed the second figure to be correct. Although the apparent prevalence of tonsillar ectopia on sagittal MRI in the study was 20%, it was explained by partial volume averaging and was therefore spurious. Secondly, in over 50% of cases with true cerebellar ectopia, the medulla oblongata is also elongated. Indeed, a linear relation has been shown between the presence of medullary elongation and severity of the descent of the cerebellar tonsils, which results in the obex often lying in the cervical canal, not the cranial cavity, and actually lying below the tonsils in about half of such cases. These anatomical facts are usually ignored in hydromechanical explanations of syringomyelia and pose appreciable difficulties. Finally, it now seems certain that the Chiari I lesion is an acquired deformation of the rhombencephalon, and not a congenital malformation at all. Serial MRI examinations have clearly shown the development of typical Chiari I deformities postnatally, the cause apparently being a lower rate of growth of the basicranium relative to the cerebellum in the first two years of life.

**Dynamics and clinical aspects**

On air myelography, an important diagnostic observation was whether an enlarged cervical cord collapsed in the head up position. Syringomyelia was collapsing, cord tumours or non-fluent cysts were not. Similar findings could be made on water soluble myelography, but the change in cord size was smaller and in the opposite direction, opacified CSF being denser than cyst fluid. Serial MRI has, however, sometimes revealed large fluctuations in cord size with no intervention whatsoever, and not associated with any change in clinical status.

There are numerous surgical strategies for collapsing a distended syrinx, the commonest being foramen magnum decompression and syringoperitoneal shunting, either of which will collapse 70%-80%. The hypothesis of Williams holds that CSF enters the cord from above, due to intermittent pathological raising of intracranial over intraspinal pressure. This has been challenged by new intraoperative measurements indicating higher intraspinal pressures, lending support to alternative hypotheses, which propose that raised intraspinal CSF pressure forces CSF into the cord via the dorsal root entry zone, or Virchow-Robin spaces. Park et al treated a small series of patients with Chiari associated syringomyelia by lumbo-peritoneal shunt-
Imaging of the spinal cord

The second. In chronic compression, although blood flow reduction is maximal in the anterior columns in contact with the compressive agent, paradoxically it is the lateral and the anterior parts of the posterior columns that show pathological change. In the brain, diffuse axonal injury is the result of shearing forces generated by rotatory acceleration. Similar forces are generated in the spinal cord, where the pia mater is restrained more than the rest of the cord structure by the duretate ligaments and spinal roots, generating shearing forces maximal in the lateral and dorsal columns. In the brain, very severe rotatory acceleration causes diffuse vascular injury, which is characterised by haemorrhagic damage in the basal ganglia, similar to the grey matter damage seen in the spinal cord. Because the white matter changes generally appear more suggestive of vascular insufficiency than diffuse axonal injury, however, most workers currently believe that progressive cord damage is due to repeated episodes of momentary arrest of the microcirculation. The resulting changes are maximal in the vascular watershed area and tend to lead to cavitation, especially in the ventral parts of the posterior columns; this is readily shown by MRI.

Imaging

Only MRI consistently shows the changes in the spinal cord that result from compression. They are best shown on T2 weighted images. The distribution is usually characteristic, consisting of diffuse signal change at the site of maximal compression, with variable extension to the central part of the cord, often bilateral, and resulting in an appearance reminiscent of snakes' eyes. These changes are shown on T1 weighted images only when the damage is particularly severe, and consist of low signal; when present they are a sign of poor prognosis. This is not so for signal changes on T2 weighted images, which often disappear completely after operative decompression, but persist when operative outcome is poor. The pathological substrate for reversible MRI changes is not known, but it is often assumed to be oedema. The size of the spinal cord at the site of compression is also of prognostic importance. Several studies have shown that when the cord is reduced in size by more than about 50%-60%, operative outcome is poor. This applies only to compression in cervical spondylosis and subluxation. The cord tolerates far greater compression from benign tumours such as meningiomas and schwannomas and functional recovery remains likely after decompression even when the cord is severely compressed.

In a recent computed myelographic study of 56 patients with spondylotic myelopathy who had a poor operative outcome, an alternative cause for the myelopathy (usually multiple sclerosis) was established in only 14-3%. The spinal cord was reduced in size by 60% or more at the site of previous compression in only 26-8%, and only 15-6% had evidence of cord necrosis. In 57-1%, operation failed to decompress the spinal canal. Another study by the same workers has put paid to the idea that osteophytes usually regress or disappear after interbody fusion: some osteophytes were as large as they had been up to eight years after the preoperative computed myelogram, and in no case did any measurable regression occur.

Clinicoradiological approach to cord compression in spondylosis and subluxations in the cervical spine

In patients with suspected compressive myelopathy, osteophytes, disc protrusions, or subluxation are irrelevant when the spinal cord is normal or only mildly flattened. Cord deformation of up to about 40% is also most likely to be irrelevant, unless appropriate signal change is present in the cord on MRI at the site of compression. Usually, however, signal change is present only in severely compressed cords.

Cord deformation of over 50% is likely to be relevant, but this is also the point at which clinical recovery from decompression is becoming less likely. Therefore, it seems appropriate to consider surgery in asymptomatic or mildly affected children and young adults as a prophylactic measure when cord compression is approaching 50%. In most patients, however, with cervical spondylosis, such operations will be less appropriate because of age and low expectation of deterioration within the relevant time frame.

Acute spinal cord injury

It is now established in animal models that the extent of signal change shown in the spinal cord on MRI is related to the severity of injury, and clinical studies have also shown a general association between the extent of signal change on MRI and functional outcome. Mild or transient loss of function after spinal injury is not usually accompanied by signal change in the cord on MRI. In more severe injury, evidence of haematomyelia is present on MRI in only about 50%. Cord swelling is mild and not always present even within seven days of injury, and ongoing cord compression is usually absent.

Progression from an acute injury to localised cystic myelopathy has been followed by serial imaging, most cysts being asymptomatic. There is good evidence that cysts result from colliquative necrosis, and extension of spindle shaped cores of ectopic necrotic tissue along the bases of the dorsal horns into adjacent uninjured parts of the spinal cord is visible on MRI. These necrotic cores are likely to be the basis of the elongated cavities that may occur within a few weeks of injury, some of which distend and propagate and become associated with a progressive ascending myelopathy. Progressive post-traumatic myelopathy can also occur in the absence of cavitation, and is associated with ascending central necrosis in the spinal cord, manifesting as signal changes on T2 and T1 weighted MR imaging.
images and abnormal accumulation of contrast medium on computed myelography. Other abnormalities such as adhesions, and occasionally cord compression, are found in some cases; some workers consider these to be the cause of progressive cord damage.

Spinal cord injuries in children differ in some ways from those in adults. Children may have extensive cord contusion or infarction with minor, remote, or no spinal fracture. Any signal change found on MRI is usually followed by considerable persistent functional loss.

Vertebral artery injury has recently been reported as occurring in nearly 46% of cases of midcervical fracture dislocation. It is notable, however, that spinal cord infarction due to vascular injury from subluxation, trauma, or cervical spondylosis and related conditions is exceptionally rare, and its documentation is confined to only two or three case reports over the past 30 years. Extensive and severe adhesive arachnoiditis and superficial siderosis are also described as rare, late complications.

Vascular lesions of the spinal cord
ANATOMY AND PHYSIOLOGY
The blood supply of the spinal cord has recently been reviewed in detail by Lasjaunias and Berenstein. The anterior spinal artery supplies a centrifugal arterial system and radial arteries from the vascular network on the surface of the cord form a centripetal system. A watershed zone between these systems has been defined consisting of the inner 25% of the white matter and the outer edge of the grey matter, excluding the posterior 50% or more of the posterior horns. Regional blood flow in cord white matter of primates has been estimated to be as low as 10 ml/100 g/min and in grey matter as 58 ml/100 g/min, which is only about half cerebral blood flow measured by similar techniques. The bases of the dorsal horns seem the most vulnerable regions within the cord to ischaemia or hypoxaemia.

Two main groups of veins drain the spinal cord. The central veins, collecting from both halves and central parts of the cord, and the radial veins from capillary plexuses at the periphery of both grey and white matter. A coronal plexus of veins on the surface of the spinal cord forms a longitudinal network which drains out of the spinal canal along the medullary veins that accompany the spinal roots at varying intervals. These veins are narrowed as they traverse the dura mater, the narrowings perhaps functioning as weak antireflux valves. Although gravity favours inferior venous drainage, in the cervical region cranial venous anastomoses seem of particular importance. High cervical obstruction has been shown to cause venous congestion and stagnant hypoxia in the central parts of the spinal cord in the cervical enlargement.

SPINAL CORD INFARCTION
The MRI appearances have been described in many cases. The commonest change has been diffuse signal increase on T2 weighted images, most often involving the lower thoracic region. Cord swelling has been mild or absent even in the acute phase. Central haemorrhage has been noted. In some cases only the ventral part of the cord has been involved, either limited in extent, confined to grey matter, or more diffusely in both grey and white matter. Diffuse contrast enhancement may be seen after intravenous gadolinium in patients examined 10–21 days after onset, but not earlier or later. An association with infarction in adjacent vertebral bodies has been noted.

Venous infarction of the cord has been reported less often, and in two recent cases the MRI abnormality consisted of unilateral signal change. One was confirmed by necropsy to be thrombosis of the posterosleral pial vein complex, and the other was speculated to be due to thrombosis in these veins induced by YAG laser during removal of an intradural neurinoma at C2. Serial MRI in the second case showed diffuse cord swelling and signal change from C1 to C3, which reduced within one month to a circumscribed area of signal change involving nearly all the lateral half of the spinal cord, confined to the site of surgery.

SPINAL VASCULAR MALFORMATIONS
Dural arteriovenous fistula
Spinal vascular malformations used to be classified according to the extent of the abnormal intradural vessels, which was a descriptive rather than a functional approach. In the early years of spinal angiography, surgeons believed that most of the vessels were arteries, as did many reputable neuropathologists until very recently. The careful observations of Kendall and Logue revolutionised thinking about these lesions, however, and now form the basis of the modern functional classification. Most spinal arteriovenous malformations are dural arteriovenous fistulae, the enlarged intradural vessels being veins not arteries. The fistula is located in the dura mater close to the nerve roots, usually in the thoracic region in older patients, but can occur in the lumbosacral theca or in the dura mater around the foramen magnum or the posterior cranial fossa. Slow, aberrant venous drainage is an important feature, and is presumed to be due to thrombosis of radicular veins. Their precise anatomy requires spinal angiography for elucidation. Treatment is often straightforward, by operative or endovascular occlusion of the fistula. Haemodynamic improvement does not always occur, however, because the thrombotic aspect of the disease may remain, with impaired venous drainage of the spinal cord. Complications of dural fistulae include intramedullary haemorrhage, cord atrophy, and cavitation in the cord, usually above the fistula.

High resolution MRI should detect most clinically relevant arterialised veins, but overdiagnosis is possible. Conventional
myelography probably remains the most sensitive and specific technique for their detection; however, enlarged or conspicuous intradural veins that drain normally, even if filled by a fistula, are not usually associated with clinical myelopathy.99 Virtually all patients with clinical myelopathy have signal changes in the lower part of the spinal cord on MRI, usually surrounded by a small rim of apparently unaltered cord tissue. The signal change often disappears partially or completely when the fistula is successfully closed and symptoms remit, and may reappear if the fistula reopens. There may be patchy enhancement of spinal cord substance after intravenous gadolinium.98,100

The site of the fistula can be detected reliably by susceptibility contrast weighted dynamic MRI when the intradural veins are large enough to be shown easily. A series of heavily susceptibility weighted fast images is acquired in the midsagittal plane after an intravenous bolus of gadolinium, and the point at which signal nulling first appears in the intradural veins indicates the level of the fistula.23 Spinal angiography can then be directed at this level, greatly speeding up the diagnostic and therapeutic process.

**Intramedullary arteriovenous malformations**

Intramedullary arteriovenous malformations may be either a nidus, or a direct arteriovenous fistula, located within cord substance or on the pia mater. Fistulae are more common in children. The nidus is often visible on MRI as a focal expansion of the cord closely associated with serpiginous signal voids indicating the draining veins and sometimes enlarged arteries. Successful endovascular treatment may be feasible, even for intramedullary lesions, but multiple sessions may be required, obliteration is often incomplete, and the recurrence rate is high.101

**Cavernomas and capillary angiomas**

Cavernomas and capillary angiomas are less common in the cord than in the brain. On MRI, they appear usually as localised expansions of the spinal cord, with sharply circumscribed signal changes that cannot be distinguished from small intramedullary haemorrhages. Intravenous gadolinium enhancement may demonstrate otherwise invisible lesions in rare cases of multiple capillary haemangiomas.103

**Neoplastic and Inflammatory intramedullary processes**

**NEOPLASTIC PROCESSES**

Astrocytomas and ependymomas occur about equally in the spinal cord itself, but ependymomas are much more common in the filum terminale. Extramedullary ependymomas occur occasionally in the extradural part of the filum, involving the sacrum.104 Glioblastomas are rare in all ages, as are oligodendrogliomas. Isolated reports are to be found of subependymomas of the cervical cord and gangliogliomas of cervicothoracic cord and filum terminale. Metastases are not uncommon. Primary lymphoma affecting only the spinal cord has now been reported several times. Reports of exceptionally rare neoplasms arising within or directly involving the cord have included melanoma; intramedullary neurofibromas arising from Schwann cells in nerves encasing blood vessels of the cord; intramedullary teratoma, associated with precocious puberty; primitive neuroectodermal tumour; mesenchymal chondrosarcoma without dural attachment, and paraganglioma of the filum terminale.

Finally, there is the relatively common spinal capillary haemangioblastoma, and its well known association with Von Hippel-Lindau disease. Screening of all family members with abdominal CT and spinal MRI with gadolinium enhancement has been recommended by several workers because 40% of affected patients may be asymptomatic at the time of screening.

On clinical imaging, the hallmark of intramedullary neoplasms is expansion of the spinal cord, usually greater than in inflammatory conditions. Lobulation, or eccentric enlargement is extremely suggestive. Both astrocytomas and especially ependymomas can appear as very well circumscribed signal change on MRI. Sometimes circumscribed lesions appear etched out by a salient low signal pseudocapsule, around the entire circumference or capping the cranial and caudal extremities, consisting of dense gliosis or haemosiderin staining; this is more frequent with ependymomas.59 Enhancement after intravenous gadolinium is usual, unlike benign intracranial gliomas, which usually do not enhance. Enhancement is patchy, and does not reliably indicate all neoplastic areas.

Haemangioblastomas and metastases usually have a different appearance. They are well defined, and enhance strongly after intravenous gadolinium. About 50% of haemangioblastomas are associated with enlarged intrathecal veins, visible on all types of imaging including MRI. Spinal cord oedema is common with metastases and shows up well in white matter with MRI11; it may be difficult to distinguish from oedema and cavitation.

Three types of cyst occur in association with intramedullary neoplasms, and about 70% will have at least one type: intratumoral cysts, the walls containing or consisting of neoplastic tissue; capping cysts, cone shaped cavities extending for one or two spinal segments into uninfiltreated cord cranial and caudal to the tumour; and syringomyelia, undistinguishable from other causes remote from the tumour.50 Even on MRI and intraoperative sonography, it can be difficult to distinguish some cystic from solid or necrotic tissue.

**INFLAMMATORY PROCESSES**

Until the era of MRI, imaging was usually negative in these conditions, but virtually all inflammatory processes produce changes in cord substance detectable by MRI. Unfortunately, they all look alike and most have been confused with neoplasia. Diagnosis
usually depends on clinical evolution, laboratory tests, or even cord biopsy, and in many cases the diagnosis remains uncertain.

**Multiple sclerosis**

The primary demyelinations, which include acute disseminated encephalomyelitis, present a spectrum of stage dependent changes, and the stages follow a roughly predictable time course that can be helpful in establishing a diagnosis.

**Stage 1: perivenous inflammation and oedema**—On MRI the cord may show mild fusiform enlargement if the lesion is large enough, with poorly defined signal change throughout the involved area, either diffuse or sparing the cord periphery. Clinical dysfunction is at its peak during this phase. Patchy or diffuse enhancement occurs after intravenous gadolinium within the area of signal change, but not coextensive with it. Similarity with cord glioma is particularly close at this stage, which lasts two to eight weeks.

**Stage 2: demyelination and glial proliferation**—Once MRI cord swelling has subsided, a smaller, more circumscribed area of signal change is evident, and enhancement no longer occurs after intravenous gadolinium. This is how most multiple sclerosis usually presents on MR images. Visualisation is considerably improved by the heavy T2 weighting provided by spinal FLAIR MRI. Lesions tend to involve sectors of the cord white matter extending to the periphery of the cord and are best shown on cross sectional T2 weighted images. The posterior columns and the posterior parts of the lateral columns are most commonly involved. The lesions are usually unilateral, or extend across the midline; they do not have the appearance of snakes’ eyes. These features should distinguish plaques of demyelination arising in the cervical cord near sites of spondylotic compression, from cord damage due to the compression alone.

**Stage 3: atrophy**—The spinal cord is small or focally or diffusely flattened. In rare cases, it becomes cavitated. Nearly all multiple sclerosis lesions eventually progress to stage 3. Acute disseminated encephalomyelitis lesions usually arrest before this stage, often not progressing beyond stage 1, and most regress completely. The prognostic significance of brain lesions, which are found at presentation in about 60% of patients with clinically isolated cord syndromes, has been reviewed recently.

**Sarcoidosis**

Involvement of the spinal cord is much less frequent than that of the brain or peripheral nerves in established cases. The appearances on myelography and MRI can be dramatic, although only a few cases have been fully described. The cord may show pronounced and extensive fusiform or irregular expansion, with variable signal changes on T1 and poorly circumscribed high signal on T2 weighted images. Patchy, non-uniform enhancement usually occurs after intravenous gadolinium and may persist for months. The solid enhancing areas have been shown to consist of astrocytic gliosis in which are embedded typical sarcoid granulomas. One operated case was also found to have extensive cystic change involving almost the entire cord, the cyst containing xanthochromic fluid. Milder forms, indistinguishable from focal multiple sclerosis lesions, have also been documented. The cord is usually involved along with the brain. Lexa and Grossman described cord involvement in three of 24 established cases of neurosarcoidosis. The association of changes in periventricular or peripheral white matter in the brain, and leptomeningeal enhancement after intravenous gadolinium, is particularly suggestive of sarcoidosis. Intravenous gadolinium is definitely helpful in identifying meningeval disease and locating additional lesions, which may clinch a difficult diagnosis. Rapid reduction in contrast enhancement, accompanied by clinical improvement, was seen in 90% of patients in response to steroid treatment.

**Spinal tuberculosis**

Spinal tuberculosis has a range of involvement similar to that of sarcoidosis. Meningeal fibrosis with chronic cavitative myelopathy is more common, especially in countries where tuberculosis has a high prevalence, such as in India, and an MR appearance consisting of multiple superficial enhancing lesions after intravenous gadolinium is probably seen more often. The diagnosis should be made from the CSF. Response to antituberculous treatment is variable, as in the brain, and may be preceded by a period of apparent worsening of the appearances.

**Intramedullary abscess (pyomyelia)**

Pyomyelia may occur from haematogenous dissemination, but is exceptionally rare. More often there is an underlying abnormality, such as a dermal sinus. A peripherally enhancing liquefying mass in a swollen oedematous cord is shown by MRI.

**Acute varicella myelopathy**

Herpes zoster can present with neurological disability before the onset of the cutaneous rash, usually consisting of unilateral limb weakness with or without long tract signs. Magnetic resonance imaging has shown mild enlargement of the spinal cord, with diffuse signal change in the ipsilateral posterolateral portion and coextensive enhancement after intravenous gadolinium. Three or four segments are involved, a little more extensive than the dermatome distribution of the cutaneous rash when it appears. Only partial resolution may follow, with residual signal change in the cord and persistence of some dysfunction. The condition can occur in fit patients as well as those who are immunosuppressed. It is considered to be due to direct involvement of the spinal cord by the virus.

**Tropical spastic paraparesis**

Tropical spastic paraparesis is a progressive vacuolar leukomyelopathy showing a strong
association with human T cell lymphotrophic virus type 1 HTLV I. The clinical course is relentlessly progressive. The thoracic region is usually involved. Extensive patchy signal change has been shown in the dorsolateral part of the spinal cord, with patchy, sometimes superficial, enhancement after intravenous gadolinium.

Listeria meningoencephalomyelitis

Listeria monocytogenes produces an encephalomyelitis characterised by multiple micro-abscesses. Mass lesion case forms, mimulating tumours. Extensive brain stem and spinal cord involvement has been reported several times. A case presenting as an isolated abscess in the cervical spinal cord was described recently, showing the MRI features of an abscess.

Lyme disease

Lyme disease may cause an acute transverse myelitis with extensive cord involvement, often associated with involvement of the peripheral nerves. Demaerel et al recently described a case involving just the spinal meninges, and we have encountered a similar case. In both, only postgadolinium MRI was abnormal, showing pronounced, diffuse enhancement of the pia mater of the brain stem and entire spinal cord. In our case, MRI of the head a few hours after the spinal examination showed that the gadolinium had diffused into the CSF producing a positive contrast cisternogram.

Granulomatous angitis of the spinal cord

Granulomatous angitis is a condition characterised by granulomata involving vascular walls, disseminated through the meninges and neural tissue, which only rarely involves the spinal cord. A case with extensive signal change throughout the spinal cord, showing no enhancement after intravenous gadolinium, has been reported, and another otherwise similar case, which showed extensive mainly superficial enhancement after gadolinium suggestive of metastatic disease ensheathing the cord.

Congenital abnormalities of the spinal cord

Congenital abnormalities of the spinal cord have been extensively reviewed by Naïdich et al, to whom the interested reader is referred for details. Some represent disorders of neurulation of the neural plate and disjunction of the neuroectoderm from the ectoderm; and these include meningo(myelo)cele and lipomas of the spinal cord. In the first, the un-neurulated neural plate (placode) remains part of the integument. In the second, a localised region of the neural tube has failed to neurulate before disjunction occurred and mesenchyme contacting the exposed dorsal surface of the neural plate has differentiated into adipose tissue; disjunction usually was complete and the overlying dura mater is intact, to create an apparently intramedullary intradural lipoma. Spinal roots emerge from the ventral surface of the un-neurulated neural placode in both conditions.

The commonest malformations involve the more caudal part of the neural tube, most of which forms by canalisation of the caudal cell mass that develops in the tail fold of the embryo. When the tail fold disappears, this part of the cord normally undergoes regressive differentiation to form the filum terminale. A useful descriptive term for this group of conditions is lipomyelomeningo-dysplasia to emphasise the elements usually present to some degree in all. The spinal cord or thickened filum extends down, usually to the sacral segments, and blends with a lipoma that extends through a dural defect and neural arch defect of variable size and length to blend with subcutaneous fat. The site of blending with the dura mater and its extent, usually referred to as “tethering”, is variable, as is the size and distribution of the lipoma and degree of meningeal ectasia. The spinal cord does not expand into a normal lumbar enlargement and the conus medullaris usually lies above or below L3, its position sometimes being difficult to define.

Another group of conditions seems to be due to much more focal, even punctate, failures of disjunction. A dorsal dermal sinus extends from skin dimple through or between neural arches to the dura, and very occasionally intradurally; about 20% of spinal dermoids and epidermoids are connected to a dorsal dermal sinus. The neuroectoderm, and ectoderm are normally briefly connected in embryonic life, via the neurenteric canal or adhesion. Persistence of the normal adhesion, or aberrant adhesions at other levels, can result in a connection from foreut to spinal canal, along which neurenteric cysts may form. Intradural spinal neurenteric cysts were well reviewed recently by Brooks et al, and present a reasonably characteristic appearance on MRI. Persistence of a communication with the skin of the back is a dorsal enteric fistula.

Diastematomyelia is a relatively common anomaly often also considered to be due to aberrant neuroentodermal adhesions. Over a variable number of segments the spinal cord develops as two, usually unequal hemicords, not duplications, although often there are two central canals, median sulci, and anterior spinal arteries. Sometimes this division takes the form only of a deep cleft, but usually the hemicords are entirely separate. In over 50% of cases, both are enclosed in a common dural tube; the rest are associated with splitting of the dura mater also, and a bony spur arising from malformed thickened laminae then often penetrates between the dural tubes. The spinal cord can be affected at any level, or rarely the filum terminale or medulla oblongata.

Finally, excessive regressive differentiation of the tail fold and caudal cell mass can lead to varying degrees of sacral and sacrolumbar agenesis, often referred to as the
caudal regression syndromes. The spinal cord is abnormally short, the conus lying in the throracic region at a variable level.

Many of these often dramatic anomalies are asymptomatic, and remain so throughout life. Much recent medical literature is still concerned with their diagnosis, which is now easy, and often seems to exaggerate the importance of timely surgical intervention. This is especially true in the concept of tethering. It has been documented recently that the filum terminale is thicker than 2 mm and filled with fat in 4% of normal patients, and that the conus medullaris lies at the level of the lower part of L2 in about 2% of the normal population. Some workers have indicated that the conus may be tethered, but normal in position, and others that functional MRI or sonography may demonstrate tethering in the absence of any morphological abnormalities. I remain aloof from such opinions at the present time. In cases with progressive disability due to the lesion, structural imaging can be important in demonstrating normal cord tissue giving rise to spinthalamic and vestibulo-spinal nerves and in suggesting an operative goal such as debulking of a lipoma or the drainage of a cyst. High resolution MRI has replaced the need for myelography and CT in preparative assessment of such cases.

Imaging of the spinal cord


**NEUROLOGY IN LITERATURE**

### Disorders of higher cortical function

I have written before of Darius Clayhanger's dressing apraxia. The account was based on Bennett's father. The underlying pathological process is not clear from Bennett's journals although I have suggested that it is possible that the condition was a rare form of Pick's disease. Bennett returns briefly to the problem in *These twain*. Most of the other extracts are concerned with memory failure either as part of senescence, or as part of a specific demetting illness. Mr Candy, in *The moonstone*, develops a memory disorder after a flu-like illness, conceivably, therefore, the sequela of an encephalitic illness. Proust's musings, expressed in his typically convoluted sentences, belong more in the realm of philosophy than neurology. It has been suggested elsewhere that Mrs Gradgrind's curious inability to relate her pain to her own body is part of a parietal disorder, although there is nothing in the novel to support that interpretation.

**Jonathan Swift, 1726, Gulliver's travels**

At ninety they lose their teeth and hair; they have at that age no distinction of taste, but eat and drink whatever they can get, without relish or appetite. The diseases they were subject to, still continue without increasing or diminishing. In talking, they forget the common appellation of things, and the names of persons, even of those who are their nearest friends and relations. For the same reason, they can never amuse themselves with reading, because their memory will not serve to carry them from the beginning of a sentence to the end; and by this defect, they are deprived of the only entertainment whereof they might otherwise be capable.

**Charles Dickens, 1854, *Hard times***

"I think there's a pain somewhere in the room," said Mrs Gradgrind, "but I couldn't positively say that I have got it."

**George Eliot, 1863, *Romola***

He was not mad; for he carried within him the piteous stamp of sanity, the clear consciousness of shattered faculties; he measured his own feebleness. Would any believe that he had ever had a mind filled with rare knowledge, busy with close thoughts, ready with various speech? It had all slipped away from him—that laboriously gathered store. . . but he found, to his acute distress, that of the new details he learned he could only retain a few, and those only by continual repetition; and he began to be afraid of listening to any new discourse, lest it should obliterate what he was already striving to remember. . . .

Old men's eyes are like old men's memories; they are strongest for things a long way off.

**Willie Collins, 1868, *The moonstone***

Here, he got on glibbily enough. Trumpery little scandals and quarrels in the town, some of them as much as a month old, appeared to recur to his memory readily. He chattered on, with something of the smooth gossiping fluency of former times. But there were moments, even in the full flow of his talkativeness, when he suddenly hesitated—looked at me for a moment with the vacant inquiry once more in his eyes—controlled himself—and went on again.

**FYodor Dostoevsky, 1869, *The idiot***

The General talked for ten minutes, heatedly and rapidly, as though too engrossed for time to express the thoughts that crowded in his head; towards the end, tears glistened in his eyes. And yet it was only sentences without beginning or end, unexpected words and unexpected ideas, rapidly and unexpectedly bursting forth and stumbling over one another.

**Arnold Bennett, 1910, *Clayhanger***

For many months now he had helped Darius to dress, when he came up from the shop for breakfast, and to undress in the evening. It was not that his father lacked the strength, but he would somehow lose himself in the maze of his garments, and apparently he could never remember the proper order of donning or doffing them. Sometimes he would ask, "Am I dressing or undressing?" And he would be capable of so involving himself in a shirt, if Edwin were not there to direct, that much patience was needed for his extrication. His misapprehensions and mistakes frequently reached the grotesque. As habit threw them and more intimately together, the trusting dependence of Darius on Edwin increased. At morning and evening the expression of that intensely mournful visage seemed to be saying as its gaze met Edwin's, "here is the one clear-sighted, powerful being who can guide me through this complex and frightful problem of my clothes." A suit, for Darius, had become as intricate as a quadratic equation.

**Arnold Bennett, 1916, *These twain***

. . . but it had witnessed hundreds of monotonous tragic meals at which the progress of his father's mental malady and the approach of his death could be measured by the old man's increasing disability to distinguish between his knife and his fork.

**Marcel Proust, 1919, *Remembrance of things past: within a budding grove***

For our memory, relatively to the complexity of the impressions which it has to face while we are listening, is infinitesimal, as brief as the memory of a man who in his sleep thinks of a thousand things and at once forgets them, or as that of a man in his second childhood who cannot recall a minute afterwards what one has first said to him.

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