Multiple lesions in cerebral white matter in two young adults with thoracic extramedullary tumours

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Abstract
Cranial MRI showed multiple lesions in white matter that were thought to be consistent with multiple sclerosis in two young adults presenting with symptoms of progressive myelopathy. MRI of the cervicothoracic spine around one and two years after onset showed the myelopathy to be due to mid-thoracic tumours. The tumours (an extradural meningioma and intradural neuroma) were resected with complete resolution of myelopathy in one patient but no recovery in the other. Spinal MRI (or myelography) should be performed in young patients presenting with signs of progressive myelopathy even when cranial MRI shows a picture typical of multiple sclerosis.

Multiple sclerosis ranks first among the non-traumatic causes of myelopathy in young adults.1 Although the relapsing remitting course of the disease usually distinguishes patients with multiple sclerosis related myelopathy from those with other aetiologies, the existence of a primary chronic progressive form of multiple sclerosis related myelopathy is well known.2

Spinal MRI has replaced myelography in investigating possible structural changes in patients with progressive myelopathies. In young adults such lesions are represented predominantly by intramedullary tumours, whereas extramedullary tumours are rather uncommon.3 Cranial MRI has been considered sufficient to diagnose multiple sclerosis related myelopathy by detecting multiple lesions in white matter in patients with negative results in investigations for spinal disease.2,4,6

We report the cases of progressive myelopathy in two young adults in which a diagnosis of multiple sclerosis related myelopathy—supported by multiple focal lesions in white matter on cranial MRI—was subsequently shown to be incorrect on spinal MRI.

Case reports
Case 1
A 20 year old man presented in April 1989 with increasing difficulty in running and a sensation of tight legs. A neurological examination at another hospital showed a mild spastic paraparesis, nystagmus on extreme lateral gaze, and decreased vibrational sense in his right leg. He had cranial MRI (0.5 T), which detected multiple tiny lesions in the subcortical and periventricular white matter (figure 1); these were thought to be consistent with multiple sclerosis. Visual evoked potentials were unremarkable. Despite steroid treatment the weakness in his lower limbs worsened, and in November 1989 he noted urinary urgency and an electrical feeling down the spine when he flexed his head. In January 1990 neurological examination showed severe paraparesis with Babinski’s sign bilaterally, clonus of the right ankle, absent abdominal reflexes, and decreased pain sensation to the lower limbs. Somatosensory evoked potentials showed a slight delay in the latency of the response after stimulation of the left median nerve and complete absence of the evoked response after stimulation of his lower limbs.
In March 1990 he was almost unable to walk, even with help; he had MRI of the cervicothoracic spine (0.5 T), which showed an extradural mass at T5 that substituted the epidural fat and impinged posteriorly on the cord (figure 2). Myelography with water soluble contrast medium showed complete block of the iodinated column at the T4–T5 intervertebral space, suggesting an extradural mass. Surgery was performed eight days after spinal MRI, and an extradural mass involving the right T4 root was removed. Histological examination showed a transitional meningioma. The patient showed prompt improvement of the paraparesis postoperatively, with complete recovery of function in January 1991.

Case 2

A 32 year old man had had pain in his left abdomen for about two years; it progressively extended to his left hip and leg and was thereafter accompanied by weakness of his left lower limb. During the two months before admission in September 1990 weakness extended to his right lower limb and he noted urinary urgency. Cranial MRI in August 1990 with an 0.5 T machine showed multiple small lesions in the cerebral white matter (figure 3) that were thought to be consistent with multiple sclerosis. At admission the paraparesis confined him to a wheelchair. Neurological examination showed brisk reflexes in his lower limbs, with Babinski’s sign bilaterally and a level of complete sensory deficit at T10. In addition, he presented with a non-paralytic, horizontal strabismus (present since his childhood, as proved in some of his old photographs), and a bilateral myopic choroid crescent was detected on fundoscopy. Thoracic MRI (0.5 T) showed an extramedullary lesion at T7 (figure 4). Although the intradural mass originating from the left T8 posterior route was completely excised at surgery nine days after spinal MRI, recovery of function was incomplete in his lower limbs; discharge was in November 1990. Histological examination showed a neuroma.

Discussion

The occurrence of monosymptomatic progressive spastic myelopathy of unknown aetiology is a comparatively common diagnostic dilemma in clinical practice. Based on the results of a necropsy series, Marshall in 1955 pointed out that multiple sclerosis is one of the main causes of these symptoms. In contrast to structural myelopathies, however, primary progressive demyelinating myelopathy predominantly affects middle aged people and is characterised clinically by a progressive neurological deficit over several years, an early onset of disturbances of bladder function, and rarity of radicular pain or loss of sensory deficit. A differential diagnosis based solely on clinical findings may be difficult in some cases. Strategies therefore have been developed to increase the detection of cases related to multiple sclerosis myelopathy by using paracranial tests. These include several neurophysiological investigations and cranial MRI—although the aim of detecting additional, clinically silent lesions of white matter—and specialised laboratory techniques such as agarose electrophoresis to detect characteristic inflammatory abnormalities of the cerebrospinal fluid.

Disappointingly, cerebrospinal fluid was not analysed in our two patients with progressive myelopathy; however, in both, cranial MRI showed multiple focal lesions in cerebral white matter, which made a diagnosis of multiple sclerosis related myelopathy probable. Although spinal MRI makes possible visualisation of spinal cord plaques in an appreciable number of patients presenting with demyelinating myelopathies, this possibility was initially discarded in our cases. This deferred the spinal MRI examination that detected extramedullary tumours as the cause of myelopathy and permitted the patient in case 1, with the shorter interval between onset and surgery, a complete recovery of the compressive myelopathy after removal of the mass. Focal lesions in white matter detected by cranial MRI are a non-specific finding in that they have been reported in several healthy elderly volunteers and have several underlying pathological changes, including demyelination plaques, microinfections, gliosis, and dilated perivascular spaces. Their observation in young adults is, however, uncommon and strongly supports a diagnosis of multiple sclerosis. Moreover, their detection in patients with unexplained myelopathy has been considered to be an important clue in diagnosing multiple sclerosis related myelopathy. Since we have no pathological correlation of the white matter lesions observed on cranial MRI in our cases we cannot establish their cause; even the hypothesis that they correspond to demyelination foci of multiple sclerosis, and hence that these patients had a dual pathology, cannot be ruled out. However, from a practical point of view, their interpretation as an indirect clue to the inflammatory origin of the progressive myelopathy in our cases proved to be wrong and misleading. This supports the opinion that spinal MRI or, if this is not available, myelography should be performed early in

Figure 2  Case 1. MRI of the thoracic spine. On a T1 weighted SE image (TR 360 ms, TE 29 ms) an extramedullary mass is detected within the spinal canal at the level of T5 and impinges posteriorly on the cord. The mass appears isointense to the cord and markedly hypointense to the posterior epidural fat.

Figure 3  Case 2. Multiple small foci areas of high signal intensity are identifiable within the periventricular and subcortical white matter on a T2 weighted SE image (TR 1600 ms, TE 100 ms).

Figure 4  Case 2. A mass hyperintense to the normal cord is present within the spinal canal at level T7 on a T2 weighted SE image (TR 1520 ms, TE 200 ms).
young patients with progressive myelopathy irrespective of the results of the cranial MRI.

3 Enzmann DR, DeLaPaz RL. Tumor: Enzmann DR, DeLaPaz RL, Rubin J, eds. Magnetic resonance of the spine, Morsby, St Louis, Missouri: 1990;301-422.