Subacute Progressive Ascending Myelopathy from L2 to C4 after A Burst Fracture of the Second Lumbar Vertebra

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Abstract

Subacute progressive ascending myelopathy is a rare, poorly understood neurological complication of spinal cord injury, unrelated to mechanical compression, instability, or syrinx formation at the level of injury or above. To date, there is no known treatment for this dramatic spinal cord injury complication. We present a case of subacute progressive myelopathy after lumbar spine trauma. The therapy consisted of plasmapheresis, hyperbaric oxygen, high-dose cortisol, antibiotic, and antiviral drugs. At 1 year post injury, the patient had recovered most of his lost upper-extremity function and MRI demonstrated only discrete signal intensity alterations extending to T3/4.

Keywords: Spinal cord injury; Fracture fixation; Plasmapheresis; Hyperbaric oxygenation; Necrosis; Apoptosis

Introduction

Subacute progressive ascending myelopathy (SPAM) is a rare, poorly understood neurological complication of spinal cord injury (SCI) [1-6], unrelated to mechanical compression, instability, or syrinx formation at the level of injury or above. Patients typically present with an ascending neurological deficit within 4 weeks after the initial injury. Characteristic magnetic resonance imaging (MRI) changes are cord expansion and an increased signal on T2-weighted sequences with a medullary distribution [6]. Only one case describing myelopathy ascending as many as 17 levels has been described in the literature [2]. To date, there is no known treatment for this dramatic SCI complication.

Case Presentation

A 27-year-old man sustained a complex fracture of the second lumbar vertebra with dura mater and nerve root injury after crashing into a tree in a skiing accident. Initial neurological examination showed incomplete conus/cauda syndrome. The cervical and thoracic spine was clinically and radiologically normal. The patient did not have any other injuries. A single dose of methylprednisolone 30 mg/kg was administered in the field. Decompression, removal of a bone fragment from the spinal canal, dural tear repair, and posterior surgical stabilization from L1 to L3 using USS instrumentation with placement of pedicle screws and an iliac crest bone graft were performed on the day of the injury. Postoperative computed tomography (CT) showed a misplaced screw in the right pedicle at L1 and the patient underwent revision on the next day. The immediate postsurgical period was uneventful. On posttrauma day 3, neurological examination revealed complete paraplegia at T12 right and L1 left. A T2-weighted MR image showed increased signal intensity within the center of the spinal cord from L2 to T11 as well as a hypointense lesion measuring 10x4 mm at level L1, suspect of an intramedullary hematoma. Six days posttrauma, the paraplegic level had ascended to T6/7. CT angiography ruled out aortic dissection, but was inconclusive with regard to perfusion. Starting on posttrauma day 7, temperature peaks were observed with the highest peak being 37.8 degrees centigrade. Blood cultures, chest x-ray, and urine test were negative. Leukocyte count was within the normal range and CRP was negative. Consequently, the subfebrile temperature was interpreted as resorption fever. Over the next days, the patient suffered progressive neurological deterioration. On posttrauma day 12, the paraplegic level had ascended to T6/7 and the increased T2-weighted MRI signal in the central cord region extended to T6 (Figure 1). There was also a new area of increased signal intensity at T12 and L1, suggestive of degraded blood products. On posttrauma day 13, the patient was transferred to our institution.

Figure 1: Increased signal intensity reaching T6 (posttrauma day 12).

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The neurological examination revealed a paraplegic level of T5/6, the patient also suffered from burning pain in the dermatomes T1/2 on both sides. On posttrauma day 14, MRI of the spine showed high signal on T2-weighted images extending from the conus to C6/7 (Figure 2). Cerebrospinal fluid (CSF) analysis revealed barrier disruption and a moderate, predominantly monocytic reactive pleocytosis with some scattered macrophages, and was interpreted as being consistent with necrotizing myelopathy. Hyperbaric oxygen (HBO) therapy and high-dose methylprednisolone were started. Neurological deterioration continued further with ascension of hypasthesia to the upper extremities. The combination of slowly developing quadriplegia and increased CSF protein led to the suspected diagnosis of Guillain-Barré syndrome and plasmapheresis cycles were initiated. On posttrauma day 17, spinal MRI including angiography was performed. The increased T2-weighted signal in the central cord region now reached C4 (Figure 3). T2-weighted images demonstrated heterogeneous intramedullary signal, the spinal canal was dilated at level T1-2. There was inhomogenous contrast enhancement at level T8-T11/12 after gadolinium application. The MRA showed no evidence of thrombosis, vascular anomaly, or dural arteriovenous fistula. An MRI of the neurocranium was normal. On posttrauma day 18, the patient had a complete sensory level at C5. Extensive virological, serological, and immunological screening was negative. In addition to 5 cycles of plasmapheresis, HBO therapy, and high-dose cortisol, antiviral (acyclovir) and antibiotic drugs were administered. 4 weeks post trauma, the extension of the characteristic signal alterations persisted at level C4 on MRI and the patient showed only minimal improvement of neurological symptoms. At 1 year post injury, he had recovered most of his lost upper-extremity function and MRI demonstrated only discrete signal intensity alterations extending to T3/4. The latest follow-up, at 3 years post injury, showed further clinical improvement of neurological function with sensory level at T4 and motor level at T6. The 3-year-follow-up MRI showed regression of signal alterations to T4, pronounced myelon atrophy below T4, and bone fusion at L1/2 level (Figure 4).

Discussion

The syndrome of delayed, subacute neurological deterioration by four or more segments following traumatic spinal cord injury was first described by Frankel in 1968, who identified eight patients, an estimated incidence of 1.5% of all spinal cord injuries [1]. A total of only 39 cases [1-7] of SPAM have been published to date, indicating that it is either exceedingly rare or grossly underreported. In particular, the occurrence of subacute delayed myelopathy after low spinal injury seems to be very rare. The only other case describing myelopathy ascending as many as 17 levels after low spinal injury (L1 level) was reported by Belanger [2]. Several hypotheses have been put forth to explain the origin of this syndrome [2].

The arterial hypothesis assumes that anterior spinal cord artery [8] thrombosis is a potential cause of the delayed deterioration. If this had been the case in our patient, deterioration would have occurred...
abolish without daily progression and the posterior column would have been preferentially spared. However, the pattern on axial MR imaging was central and thus did not appear to respect a classic ASA distribution.

Venous thrombosis and congestive ischemia are another possible etiological mechanism in ascending myelopathy. None of the MR images in our patient demonstrated venous stasis or engorgement of surface venous structures.

Ischemia due to hypotension has been proposed as a potential mechanism for SPAM [7]. This will, however, not explain the progressive nature of the ascending myelopathy unless there are repeated or sustained hypotensive episodes. Specifically, our patient was hemodynamically stable with no reported episodes of hypotension.

Inflammatory or autoimmune reaction of the central nervous system (CNS) is another possible etiological mechanism of SPAM. It could be argued that such a response may be triggered by a traumatic event, although a distinct association between the two conditions has never been identified. Our patient, however, was extensively tested for inflammatory and autoimmune processes and all the results were negative. No involvement of other parts of the CNS was noted, which could be expected in cases of demyelinating disease.

In our patient, there was no elevated leukocyte count or CRP value, no pyrexia, CSF analysis showed no signs of infection, and virological and serological workup was negative, virtually ruling out infective myelitis as a possible etiological factor.

SCI injury triggers a complex cascade of secondary neurodegenerative phenomena that are set on by the primary injury. These secondary events include neurogenic shock, vascular insults such as hemorrhage and ischemia-reperfusion, lipid peroxidation, inflammation, excitotoxicity, intracellular calcium increment, apoptosis, and disturbance of mitochondrial function [9]. The possible role of apoptosis in the causation and progression of SPAM has been elucidated by Al-Ghatany et al. [3]. While there is now strong morphological and biochemical evidence of the presence of apoptosis after SCI [8,10-20], even at distances remote from the area of initial injury [12,15,21] and persisting for many weeks [12,14,15], the mediators of post-SCI apoptosis are not yet well understood.

A number of pharmacological neuroprotective agents targeting one or more of the known secondary injury events have been studied. Some of them have shown benefit in experimental and even in clinical trials [9]. However, they are with the exception of methylprednisolone (MP) not ready for adoption in the management of patients. MP is commonly used in the setting of acute SCI based on the results of randomized controlled trials [22,23]. However, many authors have questioned the efficacy of MP because of its marginal benefits and criticisms of study design and interpretation [24-27]. MP’s mechanism of action has been attributed to anti-inflammatory or antioxidant properties. Interestingly, recently published findings suggest that MP may have direct neuroprotective activities, selectively inhibiting oligodendrocyte but not neuronal cell death via receptor-mediated action [28].

Conclusion
In summary, it is conceivable that the delayed neurological deterioration seen in our patient was due to uncontrolled secondary injury mechanisms including necrosis and apoptosis. Whether the ex iuvantibus treatment with high-dose MP and HBO contributed to the regression of neurologic symptoms in our patient can only be speculated. In any case, it has been reported that hyperbaric oxygen accelerates neurologic recovery after spinal cord injury by ameliorating mitochondrial dysfunction in the motor cortex and spinal cord, arresting the spread of hemorrhage, reversing hypoxia, and reducing edema [29].

References


