Ankylosing spondylitis and spinal cord injury: origin, incidence, management, and avoidance

W. Bradley Jacobs, M.D., and Michael G. Fehlings, M.D., Ph.D., F.R.C.S.C.

Department of Surgery, Division of Neurosurgery, Toronto Western Hospital, University Health Network, University of Toronto, Ontario, Canada

Ankylosing spondylitis (AS) is an inflammatory rheumatic disease that primarily affects the vertebral column and sacroiliac joints. Over time, the disease process promotes extensive remodeling of the spinal axis via ligamentous ossification, vertebral joint fusion, osteoporosis, and kyphosis. These pathological changes result in a weakened vertebral column with increased susceptibility to fractures and spinal cord injury (SCI). Spinal cord injury is often exacerbated by the highly unstable nature of vertebral column fractures in AS. A high incidence of missed fractures in the ankylosed spine as well as an increased incidence of spinal epidural hematoma also worsens the severity of SCI. Spinal cord injury in AS is a complex problem associated with high morbidity and mortality rates, which can be attributed to the severity of the injury, associated medical comorbidities, and the advanced age of most patients with AS who suffer an SCI. In this paper the authors outline the factors that increase the incidence of vertebral column fractures and SCI in AS and discuss the management of SCI in patients with AS. Primary prevention strategies for SCI in patients with AS are outlined as well. (DOI: 10.3171/FOC/2008/24/1/E12)

Key Words • ankylosing spondylitis • spinal cord injury • vertebral fracture • epidural hematoma

Abbreviations used in this paper: AS = ankylosing spondylitis; CT = computed tomography; HLA = human leukocyte antigen; MR = magnetic resonance; SCI = spinal cord injury.
characterized by enthesopathy, or inflammation of ligamentous insertion points, throughout the axial skeleton. This inflammatory process promotes ectopic bone formation within the affected ligaments. The widespread enthesopathy of chronic AS thus results in ossification of the ligaments of the spinal column and within the intervertebral discs, endplates, and apophyseal structures. This extensive ectopic bone formation leads to the formation of syndesmophytes, which span the ossified nucleus pulposus at each intervertebral disc level. Through the disease process, remodeling of the vertebral body also occurs. Square vertebral bodies form as a result of acute and chronic spondylitis, with destruction and rebuildong of their cortex and spongiosa. Accordingly, advanced AS, with its universal syndesmophytosis and squared vertebral bodies, leads to the formation of the characteristic hyperkyphotic “bamboo spine.”

Although new bone formation is central to the pathogenesis of AS, this pathological entity is also associated with osteoporosis and low bone mineral density (Fig. 1).21,26,43 This seemingly paradoxical finding is attributed to an uncoupling of the bone formation and bone resorption processes. Therefore, although ectopic bone formation occurs within the inflamed vertebral entheses, bone resorption, through increased osteoclast activity, also occurs at an unregulated rate within the vertebra and promotes weakening of the spinal column.21

Ankylosing Spondylitis Increases the Risk of Vertebral Fracture

As the spine in the patient with AS fuses through ligamentous ossification and syndesmophytosis, a rigid hyperkyphotic deformity develops. Biomechanically, the fused spine is more akin to a long bone and acts as a rigid lever that is incapable of appropriately dissipating the energy of a traumatic event. These altered spinal biomechanics combined with the brittle quality of the osteoporotic bone in patients with AS greatly increase susceptibility to vertebral column fractures, even after minor, often trivial, trauma.1,14,31,34

To further increase susceptibility to spinal fractures, the majority of patients with AS have significantly impaired mobility directly related to their rigid, kyphotic spinal deformity as well as a variable degree of peripheral joint arthritis. These factors exacerbate gait unsteadiness and thus increase susceptibility to falls. This finding has been supported in numerous case series of AS-related vertebral fractures, when falls are almost invariably the most common mechanism of fracture.1,14,31,34 Authors of numerous studies have evaluated the risk factors associated with vertebral fractures in the population with AS (reviewed by Geusens et al.21), and these factors include sex (men more than women), age, low body mass index, osteoporosis, disease duration, degree of syndesmophyte formation, peripheral joint involvement, increased restriction of spinal movement, and increased occiput-to-wall distance (as a surrogate measure of kyphosis). In a large review of vertebral fractures in patients with AS, Cooper et al.14 found an odds ratio of 7.7 for clinically significant vertebral column fractures, as compared with the rate in the general population. They further noted that the cumulative incidence of vertebral fractures appears to peak at 17% in the third decade after diagnosis.

Hyperextension was the most frequently observed mechanism of injury in a series of AS-related vertebral fractures,1,19,20,30,39 although the author of one series noted that flexion mechanisms predominated in a cohort.34 The prevalence of hyperextension injuries likely reflects the vulnerability of patients with AS to falls because of progressive kyphotic angulation and an inability to properly visualize the ground ahead while walking.34 Fracture patterns are also altered in patients with AS. Because of the extensive ankylosis and syndesmophyte formation, fractures often extend through the disc space and usually involve both the anterior and posterior elements of the vertebra, thus making them highly unstable (Fig. 2).

AS and SCI: Incidence and Origin

Retrospective case series of SCIs in patients with AS suggest that 1.5–2.0% of patients with such injuries also have AS,1,34,39 an AS prevalence that is much higher than in
Ankylosing spondylitis and spinal cord injury

Fig. 2. Sagittal reformatted CT scan revealing a C6–7 interdiscal fracture sustained after a fall from standing height. Note the highly displaced nature of the fracture and the involvement of both the anterior and posterior elements of the vertebral column.

Fig. 3. Sagittal reformatted (upper) and axial (lower) CT scans showing a C5–6 interdiscal fracture. This patient initially presented to another hospital with neck pain but no neurological symptoms after falling while intoxicated. Plain radiographs were interpreted as normal, and the patient was discharged. He returned 3 days later with progressive weakness in all 4 limbs.

the general population. Using Finnish national prevalence data, Alaranta et al. more recently determined that the incidence of SCI in patients with AS was 11.4 times greater than in the population at large. A higher incidence of cervical SCI has also been noted in the population with AS, accounting for up to 84% of all cord injuries, in comparison to a 55% incidence of cervical SCI in the general spinal cord–injured population.

The incidence of complete SCI also appears to be higher in patients with AS than in the general population. Patients with AS who sustain an SCI are older than the general spinal cord–injured population, with a mean age from 55 to 61 years in various series compared with a mean age of 37 in the spinal cord–injured population at large.

The higher incidence of SCI in AS is, of course, directly correlated with the increased incidence of vertebral column fractures in patients with AS. As detailed earlier, there are numerous reasons for the increased incidence of fracture in the axial skeleton of patients with AS. Furthermore, beyond the direct consequence of mechanical compression related to vertebral fracture, there are numerous other factors that contribute to the observed increased incidence of SCI in patients with AS.

For instance, vertebral fractures are often initially missed in patients with AS (Fig. 3). This unfortunate occurrence is likely to be multifactorial in its origin. First, as discussed earlier, in patients with AS, vertebral fractures often develop in the background of trivial trauma, and thus the physician’s index of suspicion may not be appropriately raised. Second, patients with AS commonly have both acute and chronic back pain, and the appropriate fracture diagnosis can be overlooked by attributing axial pain to normal disease activity. Third, given the highly abnormal structure of the vertebral column in patients with AS, spinal fracture diagnosis can be difficult on the basis of plain radiographs alone. This possibility is further confounded by the fact that many AS-related spinal fractures are located in the lower cervical spine and are poorly visualized on radiographs, which are difficult to interpret due to projection of the shoulder girdle. Accordingly, CT should be used to image the spinal column whenever a patient with AS presents with symptoms of new neck or back pain, no matter how minor or trivial the reported mechanism of injury.

Even when fractures are appropriately diagnosed, spinal column fractures in patients with AS are notoriously unstable. This high level of instability is directly related to the AS disease process. Ankylosing spondylitis promotes the ossification of spinal ligaments, which then also fracture as part of the injury pattern, further decreasing the structural support available to the spinal column. This instability often results in highly distracted vertebral column injuries (Fig. 2). The increased severity of osseous injury correlates with an increased incidence of SCI. Moreover, the greater instability significantly increases the risk of iatrogenic SCI during patient transportation and maneuvers aimed at reducing fracture dislocation. Accordingly, great care should be used whenever patients with AS and spinal frac-
Spinal epidural hematoma occurs due to bleeding from the epidural venous plexus and/or diploë of the pathological bone. Post-traumatic spinal epidural hematoma occurs with an incidence of 10–50\% \cite{15, 20, 34} and often has devastating neurological consequences if not recognized and treated emergently.

AS and SCI: Clinical Management

Spinal cord injury can be viewed as a two-phase process in which the sum total of neurological injury results from both primary and secondary injury mechanisms (recently reviewed by Baptise and Fehlings\cite{7}). The primary injury consists of the initial mechanical injury that occurs immediately following the traumatic event as a result of energy propagation from the deformed vertebral column to the spinal cord. Secondary injury begins in the minutes to hours following the primary injury and encompasses a complex cascade of cellular and biochemical processes. Secondary injury leads to neuronal and glial injury and death in a delayed fashion. These secondary mechanisms include vascular changes that promote ischemia, hemorrhage, and impaired autoregulation; ionic fluctuations that result in a loss of neuronal membrane potential; glutamatergic excitotoxicity; free radical production and lipid peroxidation; and the initiation of a robust inflammatory response.

Whereas cell death following the primary mechanical injury is almost exclusively necrotic, secondary injury events trigger a continuum of cell death that ranges from necrosis to apoptotic cell death. Together, these complex and interrelated secondary injury mechanisms promote a delayed cell death process in the spinal cord that ultimately worsens outcome after SCI\cite{7, 36, 38}. Because it occurs in a delayed fashion, secondary injury is a potential therapeutic target, and treatment strategies that can successfully limit secondary injury would potentially limit cell death and neurological deficit after SCI. Unfortunately, current attempts to minimize secondary injury have met with limited success\cite{7}, and the only clinical option in common practice, methylprednisolone administration\cite{8}, has limited efficacy at best.

Thus, until novel efficacious neuroprotective agents become available, the main focus of treatment of SCI in the acute phase consists of measures that aim to minimize secondary injury by providing the acutely injured spinal cord with adequate vascular perfusion and oxygenation. In this sense, the central tenets of acute SCI management are the same in both the injured patient with AS and the spinal cord–injured population at large.

The management of acute SCI should occur in an intensive care unit setting where monitoring allows early detection of hemodynamic instability, cardiac rate and rhythm disturbances, respiratory dysfunction, and hypoxemia. Treatment of acute SCI in the intensive care unit has been shown to improve neurological outcome.\cite{3} Specifically, blood pressure should be adjusted to provide a mean arterial pressure $\geq 90$ mm Hg for at least the first 48 hours postinjury and $\geq 85$ mm Hg for the 1st week postinjury. Judicious monitoring of oxygenation should also be instituted and any, even transient, hypoxia should be avoided. Despite the controversy surrounding its limited efficacy and marginal risk–benefit ratio\cite{24}, the administration of methylprednisolone remains a clinical option.\cite{4} Spinal cord–injured patients should also be supported with early adequate nutrition, and aggressive prophylaxis against deep vein thrombosis and thromboembolism should be instituted. As a minimum, low-molecular-weight heparin and compression stockings are recommended.\cite{2} Careful attention to the avoidance of decubitus ulcer formation is mandatory, as is early and aggressive treatment of infectious complications such as pneumonias and genitourinary tract infections.
and medical comorbidities compared with extrar-Cardiac com-plications are common in the patient with AS and indicates apposition of the caudal frac-ture fragment on the thoracic aorta.

**Special Considerations in the Management of SCI in AS**

As discussed earlier, spinal cord–injured patients with AS are older than the general spinal cord–injured population. In accordance with this increased age, patients with AS and SCI often have additional medical comorbidities and a poorer general level of health, independent of the AS, compared with the general spinal cord–injured patient pop-ulation. Because an advanced age and medical comorbidities are both factors that worsen clinical outcome and increase the chance of death after SCI, patients with AS as a group have a higher risk of such outcomes after SCI.

Whereas the major effects of spondyloarthropathies in general, and AS in particular, are related to the axial skeleton and, to a lesser extent, the peripheral joints, extrar-ticular manifestations can be present and greatly complic-ate patient care following acute SCI. Uveitis and inflammatory bowel disease are the most common extrar-ticular features, but cardiac, pulmonary, and renal involve-ments are possible as well. Renal function can be impaired due to chronic nonsteroidal antiinflammatory use, glo-merulonephritis, or amyloid nephropathy. Cardiac complications in patients with AS are related to conduction abnormalities and valvular heart disease. Recent evidence also suggests that atherosclerotic complications di-rectly related to impaired endothelial function in the spon-dyloarthropathies may be at least partly causative. Beyond direct cardiac effects, aortitis is a well-described manifestation of AS and can be significant, especially in the setting of displaced thoracolumbar fractures lying adja-cent to the aorta (Fig. 5). In this context, reduction maneu-ners have the potential to cause injury to the diseased and weakened aorta.

Pulmonary function can also be impaired due to pul-mmonary fibrosis, interstitial lung disease, pleural thickening, and chronic pleural effusions, many of which are clinically silent until late in the disease process or until events such as SCI. Decreased aerobic capacity and poor pul-mmonary function are common in the patient with AS and appear to be directly related to deconditioning. Note that deconditioning occurs because of the effects of spinal ky-}

**Ankylosing Spondylitis and SCI: Prevention**

Given the high incidence of SCI and the increased mor-bidity and mortality rates in patients with AS who suffer an SCI, it is critical that these patients be actively engaged in primary prevention strategies to avoid this devastating complica-tion of their disease process. Patients with AS should be encouraged to install activity aids such as handrails beside all staircases and within bathrooms, to use night lights in bedrooms and bathrooms, and to avoid loose area rugs that present a tripping risk. Excessive use of alcohol should be avoided, as should all contact sports or other high-impact physical activities. Seat belts should be worn at all times while driving, and car seat headrests should be used liberally. Clearly, the trade-off in terms of restricting certain aspects of daily living far outweighs the devastating morbidity and death that accompany SCI in patients with AS.

**Conclusions**

In summary, patients with AS have an increased rate of SCI because of their increased incidence of vertebral fractures. Moreover, they more commonly incur cervical SCIs and more frequently sustain complete injuries than the pop-ulation at large. These factors combined with a highly unstable spine that is predisposed to highly distracted in-juries and spinal epidural hematoma formation further increase the severity of SCI in this population. The man-agement of SCI in the population with AS is further com-plicated by an advanced patient age and the presence of multiple medical comorbidities. The result is very high mortality rates following SCI. To successfully manage SCI in this complex patient population, the neurosurgeon must be cognizant of all facets of this disease process to deftly avoid complications and thus provide these patients with the best possible chance of survival. Primary prevention strategies are also paramount and provide the best means of avoiding SCI in this susceptible population.
References


Address correspondence to: Michael G. Fehlings, M.D., Ph.D., F.R.C.S.C., Division of Neurosurgery, Toronto Western Hospital, 4W449–399 Bathurst Street, Toronto, Ontario, Canada, M5T 2S8. email: Michael.Fehlings@uhn.on.ca.

W. B. Jacobs and M. G. Fehlings