INTRODUCTION

Thoracic spinal stenosis may result from degenerative spondylosis, ossification of the posterior longitudinal ligament (OPLL), facet hypertrophy, or ossification of the ligamentum flavum (OLF) (1). Chronic severe myelopathy caused by thoracic spinal stenosis can be reversible with appropriate decompression. Neurological deterioration after surgery is a serious complication. Prevention or immediate recovery is important, however, not always possible. Paraplegia during medical treatment may result from either procedures close to the spinal cord, such as laminectomy, vertebroplasty, spondylodesis, and peridural anesthesia, involving the risk of mechanical damage to the spinal cord, or procedures distant from the spinal cord, such as vascular surgery, angiography, radiotherapy, bronchial artery embolization, and umbilical artery injection (2). Disturbances of the blood supply or toxic mechanisms may be responsible, too. Transient paraplegia after a decompressive laminectomy for the thoracic stenosis is unusual. This article describes such a case, including clinical presentation, management, and reviews the risk of neurological deterioration.

CASE REPORT

A 50-yr-old man experienced transient paraparesis for 3 to 5 min after an accidental extension of his back in March 2000. His legs turned back soon and became normal. He presented himself to the authors one month later. He felt back pain and paresthesia on both lower extremities. The paresthesia was more severe on the left side. Physical examination showed mild ankle clonus on both sides. Knee jerks were hyperactive. Babinski response was normal on both sides. Straight leg raising test was normal on both sides. An electromyography revealed lower lumbar radiculopathy. However, magnetic resonance imaging (MRI) of the lumbar spine revealed no pathological findings.

In October 2002, he visited again due to severe tingling sensation disturbing his sleep. Physical examination showed similar features. MRI of the thoracic spine revealed OLF at the T9, T10, and T11 (Fig. 1). Decompressive laminectomy from T9 to T11 was performed without gross neurological improvement. Two weeks after the first operation, laminoplasty from C4 to C6 and additional decompressive laminectomies of T3, T4, T6, and T8 were performed. Paraparesis developed 3 hr after the second operation, which recovered spontaneously 5 hr thereafter. CT and MRI were immediately performed, but there were no corresponding lesions. Vascular compromise of the borderlines of the arterial supply by microthrombi might be responsible for the paraparesis.
ing lesions. As a possible cause of the deterioration, we suggested distortion of the cord at the compression-decompression junction. So, emergency operation for a complete laminectomy to resolve the cord kinking was planned. However, the patient and his relatives refused the emergency operation. Surprisingly, he could move his legs spontaneously, about five hours later (eight hours after the second operation). His motor functions were improved to grade 3. Four days later, we drilled out all the laminae of the thoracic spine from T2 to T11 to prevent a possible cord kinking. Although there was mild hypesthesia, his motor function improved subsequently. He was discharged walking without any aids on the 34th hospital day. Although his symptoms were stable for several months, they started to deteriorate slowly. Four months after the last operation, his symptoms were almost the same as the pre-operative status. Now, he is on an alternative medicine without additional improvements.

DISCUSSION

Thoracic stenosis from OLF is not uncommon in Korea (3-8). Surgical treatment, particularly laminectomy, was usually successful and the outcomes were very promising. However, there is a risk of acute neurologic deterioration. The incidence of paraplegia through medical treatment was reported at 0.69% in an orthopedic university hospital (2). For thoracic spinal stenosis, acute neurologic deterioration is quite common after a posterior decompressive laminectomy. Young and Baron reported the incidence as high as 14.5% through a review of the literature (9). Although there have been a few cases of transient paraparesis after surgery (10-12), it is rare that transient paraparesis should occur after decompressive laminectomy and subsequently improve. Transient or temporary paraparesis may result from intraspinal hematomas (13, 14), acute aortic dissection (15), general anesthesia (16), or

Fig. 1. Magnetic resonance imaging of the thoracic spine reveals bilateral severe ossification of the ligamentum flavum at T9, T10, and T11.

Fig. 2. Computed tomographic scans of the cervical spine revealed ossification of the posterior longitudinal ligament from C4 to C6.
epidural anesthesia (17).

Direct trauma during operation or postoperative hematoma might be a cause of paraparesis. However, the patient could move all his extremities after the surgery. In addition we could not find any postoperative hematoma or any compressive lesions as in other authors’ experiences (9). Distortion of the spinal cord may occur at the compression-decompression junction (1). The altered cerebrospinal fluid dynamics may cause cord compression (11). However, spontaneous recovery before thorough decompressive laminectomy does not support such an assumption. There are several theories for neurologic deterioration after decompressive laminectomies, such as vascular compromise, hypotension or ischemia, direct trauma, or stretching of the neural elements. Some proposed the long duration of symptoms, multiple sites of compression (1), or the degree of preoperative thoracic stenosis in CT scan (18) as a bad prognostic factor. The surgical outcome is poor in patients whose initial symptoms had lasted for more than 2 yr, and who had additional proximal stenosis. The outcome is poor especially when the distal decompression was carried out first (1). We performed the distal decompression first, however, transient paraparesis occurred after the second proximal decompression. We do not know the exact mechanism of transient paraparesis of this patient. Anesthesia below the level of the nipple (T4) and spontaneous recovery suggested that vascular compromise of the borderlines of the arterial supply by microthrombi might be responsible for the paraparesis.

Decompressive laminectomy was usually successful for the OLF. However, the rate of complication was reported as high as 13.9% (19). The risk of acute neurologic deterioration is not low in the surgical decompression for the thoracic spinal stenosis. Surgeons should be alert to such complications. It is necessary to explain the rate of neurologic deterioration and possible complications after surgery before operative intervention, especially in those patients with symptoms of long duration, severe myelopathy, or multi-level involvement.

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