MR of Parenchymal Spinal Cord Signal Change as a Sign of Active Advancement in Clinically Progressive Posttraumatic Syringomyelia


Summary: Extensive MR signal change in the cranial spinal cord parenchyma was found to be an ancillary sign of disease advancement in three patients with clinically progressive posttraumatic syringomyelia. This cranial margin of parenchymal spinal cord T2 hyperintensity resolved after cystoperitoneal shunt placement. There was a concordant reduction or disappearance of the cyst in each instance, a halt in the progression of neurologic deficit, and some reversal of signs and symptoms.

Progressive posttraumatic cystic syringomyelia is an uncommon but potentially clinically serious complication of spinal cord injury (1). Delayed posttraumatic development of syringomyelia, once thought to occur in only 1% to 4% of cases, is now increasingly recognized as a major cause of morbidity (1). We describe the magnetic resonance (MR) findings in three patients with posttraumatic syringomyelia and progressive signs and symptoms, and describe a finding that may resolve after cystoperitoneal shunting. This observation may be of use in patient treatment.

Case Reports

Case 1

A 27-year-old woman had had a trauma to the cervical spine, resulting in a C5-6 fracture dislocation, 1 year before the first MR imaging examination. At the time of the injury, the spinal column was surgically fused anteriorly at three levels (C4-6) to correct posttraumatic spinal instability. Initially, the patient had a stable quadriplegia. During the 6 months preceding the first MR imaging study, she noted a progression of ascending signs and symptoms with increasing weakness in the left triceps muscle and paresthesias in the left side of the body and face. The MR study revealed a syringomyelia extending from the site of vertebral injury at C5-6 cranial to the level of the medulla (Fig 1A). There was an extensive margin of parenchymal hyperintensity along the cranial border of the cystic cavity on the T2-weighted acquisitions (Fig 1B). Neither cyst extension nor extensive parenchymal MR signal change could be identified caudal to the site of the primary injury to the cervical spine and cord. A narrow margin of parenchymal hyperintensity was noted along the caudal border of the cyst; however, it was confined to the area immediately surrounding the syrinx and the primary spinal cord injury at C-6. Subsequently, the patient had a cystoperitoneal shunt placed surgically at the C5-6 level, during which the shunt tip was advanced approximately 2 to 3 cm into the cystic cavity.

The patient was followed up clinically and with MR imaging at 2 weeks (Fig 1C) and 2 months after this intervention (Fig 1D and E). After shunt placement, the cyst gradually collapsed to the point of subtotal resolution. With this alteration, the cranial margin of parenchymal T2 hyperintensity also completely resolved. The narrow band of parenchymal hyperintensity caudal to the cyst did not resolve after surgical shunting. The patient's clinical status stabilized and somewhat improved in tandem with the imaging changes. At 1 year clinical follow-up, her signs and symptoms remained stable.

Case 2

A 52-year-old man had had a motor vehicle accident 15 years before the present admission, which resulted in a compression fracture of the L-1 vertebral body. At the time of presentation, he reported left-sided burning pain involving the inside of his thigh and the medial aspect of his calf and foot, which had been worsening during the past 5 years. The pain was described as constant and increased with coughing. On coughing he also experienced shooting pain down the posterior aspect of his left arm. The patient denied weakness except in his right leg, which had resulted in a limp beginning approximately 2 years earlier. He had daily enuresis, and also reported several incidents of fecal incontinence during the past few months.

The initial MR imaging examination revealed a syringomyelia extending from the site of vertebral injury at L-1 cranial to the level of T1-2 (Fig 2A–D). There was an extensive margin of hyperintensity along the upper border of the cystic cavity on the T2-weighted images extending cranial to the C-3 level (Fig 2B). This parenchymal hyperintensity was primarily located centrally in the spinal cord, although some relative sparing of the deep gray matter could be identified on axial images (Fig 2C). No cyst extension or major parenchymal MR signal change could be detected along the caudal margin of the cyst, because the posttraumatic cyst began at the terminal end of the spinal cord. A cystoperitoneal shunt was inserted into the cyst at the T5-6 level, and the tip was advanced approximately 2 to 3 cm into the cystic cavity.

Two months after shunt placement a repeat MR examination showed that the cyst had collapsed to the point of subtotal resolution. Along with this alteration, the cranial margin of parenchymal T2 hyperintensity completely resolved (Fig 2E and F). The patient's clinical status stabilized with resolution of pain and weakness. At the 14-month clinical follow-up, the patient's signs and symptoms showed no evidence of progression.

Case 3

A 72-year-old man had a history of thoracic spine trauma 2 years previously, which resulted in a T5-6 fracture dislocation.
During the 6 months preceding the initial imaging study, the patient noted a progression of ascending signs and symptoms that included pain and paraesthesias along his upper extremities. The MR examination revealed a syringomyelia extending from the site of vertebral injury at C5-6 cranial to the level of the medulla. Note little or no caudal extension of the cyst.

The patient was sequentially followed up after surgery, clinically and with MR imaging, at 2 and 6 months. After shunt placement, the cyst gradually collapsed to the point of subtotal resolution. With this alteration, the cranial margin of parenchymal T2 hyperintensity completely resolved. The patient's clinical status stabilized and somewhat improved in concert with the imaging changes. At the 18-month clinical follow-up, the patient's signs and symptoms remained stable.

**Clinically Stable Cases**

Three patients with clinically stable posttraumatic quadriplegia were also studied by MR imaging in a manner identical to the group with clinically progressive signs and symptoms. The preceding trauma had occurred 1 to 5 years prior to the MR examinations. These patients all had focal syringomyelia in the cervical spinal cord. The cysts did not extend beyond one cervical vertebral body length. In one case, the spinal cord showed no discernible pericystic T2 signal change, while in the
other two cases the T2 signal change was confined to the parenchyma of the spinal cord immediately surrounding the parenchymal cyst, not exceeding one vertebral body length.

**Discussion**

Progressive signs and symptoms related to posttraumatic syringomyelia can occur as a late sequela to spinal cord injury (2–4). The prevalence of posttraumatic syringomyelia is much higher in patients with complete quadriplegia (8%) than in those with incomplete quadriplegia (2.5%), complete paraplegia (1%), or incomplete paraplegia (2%) (4). Signs and symptoms of posttraumatic syringomyelia have been reported as early as 3 months after spinal cord trauma or as late as 32 years after injury (4).

Ascending and descending posttraumatic myelopa-
thy is thought to begin at the time of injury or soon thereafter and may involve mechanisms such as the release of destructive enzymes, free radical products, or other toxic agents (5–8). Central or dorsocentral cores of necrotic tissue may extend for several spinal segments from the site of vertebral injury at T-5 (not shown) to the level of C4-5.

Central or dorsocentral cores of necrotic tissue may extend for several spinal segments from the site of vertebral injury at T-5 (not shown) to the level of C4-5.

Conversion of necrotic spinal cord into a true cyst may take months to years (4, 7, 9). This probably accounts for the apparent continuum of noncystic to cystic myelopathy. The rate of progression of the pathologic process in individual cases is highly variable (7).

The pathogenesis of syringomyelia formation is unclear, although several theories have been proposed. Kao et al (10) have postulated that the rupture and coalescence of microcysts lead to posttraumatic syringomyelia formation. Microcyst formation in contused, compressed, and transected spinal cords results in part from the egress of fluid from damaged axons (11, 12). Other proposed mechanisms that lead to formation of posttraumatic syringomyelic cavities include resorption of blood and necrotic tissue, lysosomal autodigestion, secretion of cystic fluid by the developing cyst lining, infarction, posttraumatic reactive ependymal proliferation causing segmental closure of the central canal with resulting local distension, and passage of cerebrospinal fluid (CSF) into the injured central cord via enlarged perivascular spaces (1, 2, 10, 12–17). The formation of arachnoid adhesions may further alter the CSF dynamics so that abnormal...
entry of CSF into the spinal cord via Virchow-Robin spaces results in progressive syringomyelia (7, 18, 19).

Reports indicate that posttraumatic syringomyelia most commonly extends superiorly from the injury site; less frequently, it may also have an inferior component. Inferior extension alone is uncommon (9). It has been theorized that because there is more space in the cervical than in the thoracic spinal canal, the cervical cord can expand more easily from the underlying cystic cavitation (20, 21). An additional limitation in caudal growth of cysts may be the posttraumatic parenchymal spinal cord atrophy and gliosis at and below the injury site, resulting in a more dense, compact cord that is therefore less able to expand. For these reasons, the cyst would be hypothetically freer to extend along the line of least resistance in the cranial direction (21).

The first clinical suggestion of advancement of a posttraumatic syringomyelia is usually the onset of ascending signs or the development of new symptoms after an apparently stable period (1, 22). The progressive clinical picture varies from a new onset of minimal sensory loss to a fully developed anesthesia with motor weakness of an extremity and the development of Charcot joints (12, 14, 22). In some patients, the chief symptom is severe pain (4, 22). Although sensory impairment is troublesome initially in many cases, in later stages motor loss may become sufficient severe to warrant further radiologic investigation or active surgical intervention.

The presence of flow voids on MR images within the syringomyelic cystic cavity has been suggested by some to correlate with a greater potential for improvement after surgical shunting (22). In our cases we did not perform specific MR examinations, such as gated CSF flow studies to determine the nature of fluid dynamics within the syringal cavities. We were therefore unable to correlate cystic fluid flow characteristics with our observations of high parenchymal signal on T2-weighted MR images above the cysts.

Gebarski et al (23) categorized posttraumatic progressive myelopathy into three groups on the basis of characteristic radiological findings and their responses to therapy: 1) myelomalacia with no cystic degeneration: poor response to therapy; 2) small cyst formation: poor response to surgical therapy; and 3) large cyst formation: effective treatment by surgical shunting.

Surgical shunting of these large cysts may halt the progression of the myelopathy and reverse the clinical deterioration in many cases (9). Hyperintense T2 MR signal within the spinal cord parenchyma adjacent to syringomyelia as shown in the current group of patients may correlate with both edema formation and myelin degeneration with or without gliosis. Such MR signal change is not typically observed distant from the site of focal cord trauma; permanent parenchymal signal alteration is usually only seen immediately surrounding the primary focus of cord trauma. Furthermore, in instances in which this parenchymal signal change only extends cephalad, and is reversible, the chances are favorable that edema is the main factor: resolution of this abnormal parenchymal MR signal intensity after cystoperitoneal shunting appears to correlate with improvement in neurologic status. Although these cranial parenchymal MR signal changes were presumably progressive, together with the cranial enlargement of the cyst, our small group of patients did not have prior imaging studies with which to compare the images acquired at the time of presentation of advancing signs and symptoms. This possibility can only be assessed in future studies of patients in whom there is more complete longitudinal imaging documentation. Nevertheless, the focality of the posttraumatic syringomyelia and the lack of extensive cranial T2 MR parenchymal spinal cord signal change in the second group of clinically stable posttraumatic quadriplegic patients supports the significance of the imaging findings in the patients with clinically progressive disease. Presumably, the T2 MR parenchymal spinal cord signal alteration that was seen immediately surrounding the spinal cord cysts in two of the clinically stable patients represented static posttraumatic gliosis (ie, myelomalacia).

In summary, some enlarging posttraumatic syringomyelic cysts are observed to have a cranial margin of parenchymal spinal cord T2 hyperintensity on MR studies. Because this abnormal parenchymal MR signal intensity resolved together with advancing signs and symptoms after surgical cystoperitoneal shunting of the cystic cavity, it may represent fluid escaping from the cyst or edema caused by as yet undefined pathologic alterations of the spinal cord adjacent to the enlarging cyst. While ascending/advancing clinical signs and symptoms are the main factors to consider in patients being considered for surgical shunting, the ancillary radiologic sign described here should be taken into account when imaging patients with posttraumatic syringomyelia and evidence of clinical progression.

References


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