Arachnoiditis ossificans associated with syringomyelia: An unusual cause of myelopathy

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ABSTRACT

Objective: The pathophysiology of arachnoiditis ossificans (AO) and its association with syringomyelia remains a rare and poorly understood phenomenon. Here, we present a case of AO associated with syringomyelia, a review of literature, and a discussion of current understanding of disease pathophysiology.

Methods: A literature review was performed using MEDLINE (January 1900–May 2010) and Embase (January 1900–May 2010) to identify all English-language studies that described AO with syringomyelia. The current report was added to published cases.

Results: Over 50 cases of AO are reported in literature, of which only eight are associated with syringomyelia. The various presumptive etiologies of syrinx formation include abnormalities in blood circulation, ischemia, hydrodynamic alternations in cerebrospinal fluid (CSF) flow, tissue damage, or incidental coexistence. Changing CSF dynamics related to decreased compliance of the subarachnoid space and subsequent paracentral dissection of the spinal cord may be implicated in the disease process. Magnetic resonance imaging (MRI) scanning may identify the syrinx but fail to diagnose the calcified arachnoid. Five patients, including the current case, improved clinically following laminectomy and decompression.

Conclusions: Syringomyelia in association in AO is a rare phenomenon. A high index of suspicion is required and both MRI and computed tomography (CT) are recommended for diagnosis. The pathophysiology of syringomyelia in AO remains an area of ongoing research.

ABBREVIATIONS LIST

AO Arachnoiditis ossificans
CSF Cerebrospinal fluid
CT Computed tomography
MRI Magnetic resonance imaging

This report has received no financial support.
Case report—Arachnoiditis ossificans associated with syringomyelia: An unusual cause of myelopathy

INTRODUCTION

Calcification of the arachnoid membranes in the intrathecal space is a common finding on autopsy [1]. However, arachnoiditis ossificans (AO) is a unique and uncommon phenomenon characterized by intrathecal bony metaplasia of the arachnoid membrane [1]. This pathology develops following numerous predisposing processes including surgery, trauma, infection, myelography and subarachnoid hemorrhage [2–4]. As of yet, the pathophysiology of AO is unknown; however, it may be a result of chronic inflammation of the arachnoid [5].

Over 50 reports of AO are documented in literature; however, only eight of these are associated with syringomyelia [2–9]. The pathophysiology of syrinx development in this disease state is controversial. Here, we present a case of AO-associated syringomyelia and provide a review of literature and discussion of the current understanding of disease pathophysiology.

CASE REPORT

History

A 36-year-old male truckdriver was referred to our neurosurgical spine service with a 4-year history of progressive spastic paraparesis and gait imbalance. His past medical history was significant for cerebellar ganglioglioma requiring craniotomy and resection 17 years prior to presentation. This was complicated by CSF leakage and subsequent surgical repair. He had no upper extremity symptomatology and his bowel and bladder function was intact. Based on his history, his neurological symptoms were initially attributed to his cerebellar pathology. His symptoms however progressed to debilitating lower extremity spasticity such that he underwent several Botox injection procedures and further investigations were carried out.

On examination, the patient was myelopathic and grossly spastic in his lower extremities with increased tone bilaterally. He was unable to perform tandem walking and Romberg's test was positive. His pain and temperature sensation was intact; although proprioception in the lower extremities was impaired.

Imaging

A plain film of the thoracic spine revealed no abnormalities. An MRI scan revealed a thoracic syrinx at the T3 level as well as an intradural extramedullary lesion displacing the spinal cord at the T4–5 levels. An MRI scan revealed a thoracic syrinx at the T3 level as well as an intradural extramedullary lesion displacing the spinal cord at the T4–5 levels.

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Evidence-Based Spine-Care Journal

Volume 1/Issue 2 — 2010
level measuring 2 cm × 5 mm. There was no evidence of enhancing tumor associated with the syrinx. An intradural extramedullary mass along the left posterolateral aspect of the thecal sac was noted at T4–5. This lesion displaced the spinal cord anteriorly and was associated with abnormal signal within the spinal cord suggestive of myelomalacia or edema. Hypointense nodularity with the appearance of intradural calcification was seen posteriorly from T5–10 (Figure 1). Our initial differential diagnoses for this intradural extramedullary lesion included: meningioma, schwannoma, a drop metastasis from the previous cerebellar gangioglioma or a primary exophytic tumor. At this point, we proceeded with a T4–7 laminectomy for spinal cord decompression.

**Operative procedure**

With monitoring of motor and somatosensory evoked potentials, a laminectomy was undertaken from T4–T7 and the thecal sac was exposed. Intraoperative ultrasound was used to identify and characterize the extent of the syrinx and calcified lesion (Figure 2). As the durotomy was performed, it was apparent that there was marked arachnoiditis with arachnoid calcification extending as low as T8–9. The lesion was visualized to be a large calcified segment of arachnoid densely adherent to the dorsal surface of the cord. Meticulous and careful dissection allowed the lesion to be elevated away from the cord surface. The lesion itself was approximately 38 × 18 mm (Figure 3). The cord was noted to be pulsatile following the decompression. A duroplasty was fashioned to reconstruct the subarachnoid space. Evoked potentials were unchanged throughout the procedure.

**Outcome**

At 9-month follow up, the patient noted marked improvement in gait. He was able to ambulate without walking aids. His spasticity also improved although he continued to receive botox injections to his lower extremities.

Repeat MRI revealed a significant decrease in the size of the lesion (Figure 4). The syrinx at T3 was significantly smaller compared to preoperatively and there was near complete resolution of the edema within the spinal cord. There was also a marked reduction in mass effect and compression of the spinal cord at the decompressed levels. Persistent adhesions of the arachnoid at other levels were noted.

**DISCUSSION**

The earliest clinicopathologic association of arachnoid calcification with syringomyelia dates to an autopsy study by Schwarz in 1897 [10]. Eight cases of arachnoiditis ossificans with associated syringomyelia have been published in literature (Table 1). In all of these cases, the patients presented with spastic paraparesis. Due to the au-
In all reported cases, the syrinx localized to the thoracic cord within close proximity to the areas of AO. A thoracic laminectomy was consistently performed and durotomy and duroplasty usually achieved. The calcified arachnoid plaques were resected to variable extents due to their firm adherence to the spinal cord. In several cases, the syrinx was aspirated and a syringoperitoneal or syringopleural shunt was inserted [5, 6, 8]. Collapse of the syrinx was noted with some postoperative clinical improvement in two of these reports [5, 8]. Clinical improvement was also noted by several authors who only performed decompression and removal of ossified plaque without myelotomy or shunting of the syrinx [7, 9]. This was also observed in the current study. Our study is the first to document a decrease in the size of the syrinx on postoperative imaging following spinal cord decompression in patients with AO without myelotomy or shunt insertion.

Domenicucci and colleagues [7] classify AO into three categories based on the calcified arachnoid morphology. Type I ossifications are semicircular in appearance, type II are circumferential and type III are a honeycomb-like ossifications, which affect the cauda equina. In their reports and review, one-third of patients with type I or II were associated with syringomyelia and/or spinal cord cysts.

Syringomyelia in the setting of AO has been hypothesized to result from vessel ischemia, CSF flow alterations, or even as a coexisting incidental finding [2, 5, 6]. In a large study of 105 patients, Milhorat and colleagues identified 23 cases of focal intramedullary cavitation separated from the fourth ventricle by a syrinx-free segment of spinal cord [11]. These occurred in association with Chiari I malformations, cervical spinal stenosis, spinal arachnoiditis, basilar impression, occipital en-cephalocele and congenital cyst of the terminal ventricle. Syringomyelia was associated with arachnoiditis in 13% of these cases. The histological features of these syrinxes differed from those that communicate with the fourth ventricle or posttraumatic extracanalicular syrinxes in that these were associated extensive ependymal denudation, paracentral dissection and intracanalicular septations. Dissections occurred at the poles or thinnest segment of syrinx and occasionally ruptured through the pia to communicate with the subarachnoid space. The authors hypothesize alterations in CSF dynamics is the impetus for disease progression.

Studies of syringomyelia associated with Chiari I malformations have hypothesized that syrinx progression is a result of cerebellar tonsils, which occlude the subarachnoid space at the foramen magnum and act as a piston on the partially enclosed spinal subarachnoid space [12]. This may exaggerate cervical subarachnoid pressure waves that compress the spinal cord externally and propel the syrinx caudally with each heartbeat. Similar CSF flow changes in arachnoiditis may explain the unique histological phenomenon of paracentral dissection and account for reported cases of rapid progression of the syrinx in AO [2].

Another compelling theory relates to decreased compliance of the subarachnoid space. In animal models of syringomyelia where arachnoiditis was induced by injection of kaolin in the subarachnoid space of the upper thoracic spinal cord, it was found that subarachnoid space pressures were higher rostral to the arachnoiditis compared to caudal [13]. Furthermore, when CSF flow was studied with injection of the CSF tracer horseradish peroxidase, it was found that perivascular spaces were enlarged and the tracer stained these spaces and the central canal within 10 minutes. A reduction in the compliance of the subarachnoid space in this setting may increase the perivascular flow of CSF contributing to the formation and enlargement of syringomyelia.

Controversy in literature remains regarding the need for drainage and shunting of the syrinx. The current study is the first to report that decompression alone results in a decrease in the syrinx size on follow-up imaging with clinical improvement of the myelopathy and spasticity. Given current understanding of syrinx pathophysiology, it is the authors’ opinion that reconstruction of the subarachnoid space in an attempt to normalize CSF fluid circulation may be adequate to decrease the syrinx size. This may avoid the added risks of myelotomy and shunting. Further research to identify the pathophysiology and natural history of syringomyelia in the setting of arachnoiditis is necessary before recommendations are made to change current practice.
**CONCLUSION**

Syringomyelia associated with arachnoiditis ossificans is a rare condition, which may be due to altered CSF dynamics secondary to the ossified arachnoid. CT scans are recommended for diagnosis as MRI may lack the sensitivity to identify intrathecal ossification and calcifications. Laminectomy and decompression with or without syrinx drainage and shunting remain the standard of care. A broader understanding of the pathophysiology of syrinx formation and progression is necessary and such research is currently underway.

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**Table 1  Summary of conditions and weights used to calculate Charlson comorbidity score**

| Clinical conditions | Weights |
|---------------------|---------|
| Myocardial infarction, congestive heart failure, peripheral vascular disease, dementia, chronic lung disease, connective tissue disease, ulcer, chronic liver disease | 1 |
| Hemiplegia, moderate or severe renal disease, diabetes with complication, tumor, leukemia, lymphoma | 2 |
| Moderate or severe liver disease | 3 |
| Malignant tumor, metastasis, AIDS | 6 |

* Weights are assigned to 19 predetermined clinical conditions with the final score being the sum of the weights assigned to the conditions with scores ranging from 0–37. One point for each additional decade of life over 50-years-of-age can be added to initial score to create a single index. The higher the weight total, the greater the likelihood of mortality.

**Acknowledgements:**
The authors thank Steve Lunny for his help in formatting the figures included in the manuscript.
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The case presented is a Domenicucci type 1, representing a case with one or more surgically resectable compressions caused by a calcified arachnoid. The recommended treatment consists of resection of the calcification for decompression and duraplasty as necessary. In the absence of improvement or with progression of symptoms associated with a concurrent syringomyelia, myelotomy for cerebrospinal fluid drainage and shunting had been recommended.

The case described in the preceding text is the first one in the literature treated with decompression alone, and is very well documented. The demonstrated decrease in the syrinx size—addition to a good clinical outcome—supports the recommendation of drainage surgery to be performed only in presence of progression of symptoms.

As a final observation this case has only 9-months of follow up, which is a short time from which to draw conclusions as to eventual surgical success.

Additional references:
1. Mello LR, Bernardes CI, Feltrin Y, et al (2001) Thoracic spine arachnoid ossification with and without cord cavitation: report of three cases. J Neurosurg: 94 (1 Suppl):115–120.
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Editorial Staff Perspective

Case reports have been frequently touted as representing the 'bottom of the food chain' in the evidence pyramid. The preceding case report explicitly presents a classic indication for a meaningful case report even in this day and age of insistence on 'higher levels of evidence'. This exceedingly rare condition was luckily identified and after thorough review treated by expert surgeons armed with skill set and expertise to handle even the rarest and most unusual of cases.

We all continue to gain by case reports in these settings, especially if they go through the extra effort to summarize the past publication history in the topic in a mini metanalysis—as was done in this case. Caution should, however, be exercised in the over application of greater inferences for larger patient populations based on such case reports.

Additional references:
1. Mello LR, Bernardes CI, Feltrin Y, et al (2001) Thoracic spine arachnoid ossification with and without cord cavitation: report of three cases. J Neurosurg: 94 (1 Suppl):115–120.
2. Faure A, Khalfallah M, Perrouin-Verbe B, et al (2002) Arachnoiditis ossificans of the cauda equina Case report and review of the literature. J Neurosurg: 97 (2 Suppl):239–243.