Dorsal arachnoid webs (DAWs) and idiopathic spinal cord herniation (SCH) are uncommon abnormalities affecting the thoracic spinal cord that can result in syringomyelia and significant neurological morbidity if left untreated. Differentiating these 2 entities on the basis of clinical presentation and radiological findings remains challenging but is of vital importance in planning a surgical approach. The authors examined the differences between DAWs and idiopathic SCH on MRI and CT myelography to improve diagnostic confidence prior to surgery.

**OBJECTIVE** Dorsal arachnoid webs (DAWs) and spinal cord herniation (SCH) are uncommon abnormalities affecting the thoracic spinal cord that can result in syringomyelia and significant neurological morbidity if left untreated. Differentiating these 2 entities on the basis of clinical presentation and radiological findings remains challenging but is of vital importance in planning a surgical approach. The authors examined the differences between DAWs and idiopathic SCH on MRI and CT myelography to improve diagnostic confidence prior to surgery.

**METHODS** Review of the picture archiving and communication system (PACS) database between 2005 and 2015 identified 6 patients with DAW and 5 with SCH. Clinical data including demographic information, presenting symptoms and neurological signs, and surgical reports were collected from the electronic medical records. Ten of the 11 patients underwent MRI. CT myelography was performed in 3 patients with DAW and in 1 patient with SCH. Imaging studies were analyzed by 2 board-certified neuroradiologists for the following features: 1) location of the deformity; 2) presence or absence of cord signal abnormality or syringomyelia; 3) visible arachnoid web; 4) presence of a dural defect; 5) nature of dorsal cord indentation (abrupt “scalpel sign” vs “C”-shaped); 6) focal ventral cord kink; 7) presence of the nuclear trail sign (endplate irregularity, sclerosis, and/or disc-space calcification that could suggest a migratory path of a herniated disc); and 8) visualization of a complete plane of CSF ventral to the deformity.

**RESULTS** The scalpel sign was positive in all patients with DAW. The dorsal indentation was C-shaped in 5 of 6 patients with SCH. The ventral subarachnoid space was preserved in all patients with DAW and interrupted in cases of SCH. In no patient was a web or a dural defect identified.

**CONCLUSIONS** DAW and SCH can be reliably distinguished on imaging by scrutinizing the nature of the dorsal indentation and the integrity of the ventral subarachnoid space at the level of the cord deformity.

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**KEY WORDS** arachnoid web; cord herniation; MRI; CT myelography; congenital
syndrome. The radiological differentiation of these entities may be similar as well. In a review of 27 patients with SCH by Tekkök, a correct preoperative diagnosis was made in only 9, with SCH being confused most often with dorsal arachnoid cysts or masses. On MRI and myelography, both entities present as focal anterior displacements of the midthoracic cord with widening of the posterior subarachnoid space. The arachnoid web and the dural defect through which the cord herniates are difficult to directly visualize on imaging. Indeed, some cases of SCH were mistakenly attributed to ventral displacement of the cord by dorsally located arachnoid “cysts,” a conclusion stemming from failure to recognize that the dorsal “cyst” was simply an expanded subarachnoid space resulting from the cord being pulled into a dural defect.

We therefore sought to examine the differences between these superficially similar entities on MRI and CT myelography in a series of patients who had undergone surgical lysis or release for DAWs or SCH, respectively.

Methods

This HIPAA-compliant retrospective study was approved by the institutional review board of the University of Maryland Medical Center and the need for informed consent was waived. From a review of the picture archiving and communication system (PACS) database between 2005 and 2015, we identified 6 patients with DAWs and 5 with SCH. Clinical data including demographic information, presenting symptoms and neurological signs, and surgical reports were collected from the electronic medical records. Ten of the 11 patients underwent MRI. CT myelography was performed in 3 patients with DAW and in 1 patient with SCH. MR images were obtained on 1.5-T and 3.0-T MRI machines (Siemens). MRI sequences included standard acquisition, fast spin-echo technique with T1- and T2-weighted acquisition. Sagittal T2 sampling perfection with application-optimized contrasts using different flip-angle evolution (SPACE) sequences were acquired using the following parameters: TR 1500 msec, TE 130 msec, flip angle 150°, 1-mm slice thickness, and bandwidth 465 Hz. Axial CT myelographic images were obtained following administration of intrathecal iodinated contrast using the following parameters: 120 kV, dose modulation with variable milliamperes, and 2-mm spatial resolution with coronal and sagittal reformatted images. All diagnoses were confirmed on surgery.

The imaging studies were analyzed by 2 board-certified neuroradiologists (P.R. and A.S.), both with more than 6 years of experience in diagnostic neuroimaging. The following features were assessed: 1) location of the deformity; 2) presence or absence of cord signal abnormality or syringomyelia; 3) visible arachnoid web; 4) presence of a dural defect; 5) nature of dorsal cord indentation (abrupt [“scalpel sign” as defined by Reardon et al.] vs “C”-shaped; Fig. 2); 6) focal ventral cord kink; 7) presence of the nuclear trail sign (endplate irregularity, sclerosis, and/or disc-space calcification that could suggest a migratory path of a herniated disc); and 8) visualization of a complete plane of CSF ventral to the deformity.

Results

The age range for patients with DAWs was 18–67 years (4 males, 2 females), and 45–77 years (3 men, 2 women) for patients with SCH. DAWs were encountered between the T-3 and T-6 levels and SCHs were noted between the T-5 and T-8 levels. Symptoms and clinical findings in both groups included back pain, lower-extremity weakness and paraesthesia, erectile dysfunction, and fecal and urinary incontinence. One patient with SCH had Brown-Séquard syndrome. The frequencies of the above-described imaging findings are indicated in Table 1. An arachnoid web was not definitively seen preoperatively in any patient with surgically proven DAWs. The dorsal cord indenta-
Illustrative Cases

Case 1 (DAW)

A 56-year-old man presented with a 3-month history of episodic urinary urgency, urinary incontinence, radicular pain, and gait difficulty. MRI of the thoracic spine showed expansion of the spinal cord from T-6 to T-10 with pre-syrinx conditions as well as a sudden change in cord caliber at the T-6 level with a scalpel-like dorsal indentation. T2-weighted SPACE and subsequent CT myelographic images did not show the arachnoid web directly. However, both demonstrated a complete plane of CSF ventral to the subarachnoid space visualized in all 6 patients with DAW. This was easier to determine on T2 SPACE and CT myelographic images. In all 5 patients with SCH, this plane was found to be interrupted.

Case 2 (SCH)

A 77-year-old man presented with several years of lower-extremity weakness, numbness and paresthesia, and recent onset of urinary incontinence. On examination, decreased strength in the lower extremities and loss of sensation in the right toes and perineum was noted. MRI, originally misinterpreted as a dorsal arachnoid cyst, revealed a focal cord deformity at T-5 with a C-shaped dorsal indentation and loss of the ventral CSF plane at the level of the deformity. He underwent a T5–6 laminectomy, release of the ventral herniation, dural repair, and T4–6 fusion, with recovery of sensation and motor function (Fig. 4).

Discussion

Our findings revealed that DAW and SCH can be reliably distinguished on imaging by scrutinizing the nature of the dorsal indentation and the integrity of the ventral subarachnoid space at the level of the cord deformity. The septum posticum of Schwabé traverses the posterior spinal subarachnoid space in the sagittal plane and it is especially well developed in the thoracic spinal canal where it presumably serves to stabilize the cord. It is sometimes seen as a filling defect on supine myelograms. The septum is believed to progressively degenerate with arachnoid cysts, with webs and adhesions representing its remnants. These may compress the cord and impede CSF flow to varying degrees. Most cases of SCH are idiopathic. They may arise from remote trauma leading to disruption of the dura or in some instances from a dural rent caused by a disc protrusion11 with subsequent tamponade by the cord. Rare instances of cord herniation into congenital dural defects and of herniated cord tissue trapped between 2 layers of congenital dural duplication have also been reported. An as-yet undetermined inflammatory process leading to adhesion of the cord to the ventral dura and subsequent development of a dorsal defect arising from pressure erosion by the pulsating, adherent cord has also been proposed as a mechanism for development of SCH. The clinical presentation of both entities is similar with symptoms including back, neck, or shoulder pain; upper- and lower-extremity paresthesias and weakness; and Brown-Séquard syndrome. Symptoms may be due to a combination of direct compression of the cord and development of syringomyelia, presumably resulting from impairment of CSF flow.

Surgical treatment of DAW involves laminectomy with intradural lysis of adhesions and resection of arachnoid bands. Careful attention must be given to verify that there is satisfactory flow of CSF from the cranial and caudal ends of the dural opening. If there is residual arachnoid membrane present, at the caudal or cephalad aspects of the dural opening, we advocate suturing this arachnoid layer to the residual dura during dural closure to prevent CSF from accumulating into a potential space between the arachnoid and the dura. The primary aims of SCH repair include reduction of the herniation, restoration of the spinal cord to its normal position, and prevention of recurrence of herniation. There appear to be 3 approaches toward these ends: some advocate primary closure of the dural defect, some close the dural defect with fat or fascia, and others prefer to enlarge the dural defect. All of these require inspection of the ventral cord surface by dividing the dentate ligaments. Complications from such procedures may include anterior epidural CSF collection and superficial siderosis.
Direct visualization of the DAW remains difficult using conventional imaging techniques. A web was not confidently identified in any of the 14 cases reported by Reardon et al.\textsuperscript{12} and was identified in only 2 of 7 cases by Hakky et al.\textsuperscript{8} Improved visualization of these webs may be achieved by the addition of “myelographic” high-resolution T2-weighted MRI sequences\textsuperscript{6} such as constructive interference in steady state (CISS) or SPACE, or their equivalents (Fig. 3). Despite the use of the SPACE sequence in 4 patients in our series, a web was not definitively identified in any of these patients. The dural defect and focally herniated ventral cord parenchyma may also be difficult to identify. A dural defect was not identifiable in any of our patients and a ventral cord deformity was present in only 3 of 5 patients with SCH.

Preoperative imaging differentiation between these entities is therefore best achieved by identifying indirect signs. The 2 most reliable signs to differentiate DAW from SCH were 1) the shape of the dorsal indentation on sagittal MRI or CT myelography, and 2) the presence or absence of a plane of CSF ventral to the cord. The scalpel sign described by Reardon et al. refers to a characteristic notch.
such as focal dorsal indentation of the upper thoracic spinal cord on sagittal MRI and CT myelography, resulting in the cord resembling a scalpel with its blade pointing posteriorly.\textsuperscript{12} This arises from displacement and compression of the cord by the arachnoid band in the dorsal subarachnoid space and was present in all cases of DAW in our series. In all patients with SCH, the indentation was C-shaped, due to a short segment of the cord being pulled ventrally, distal to which it returns to its normal position (Figs. 1 and 2). Close scrutiny of the ventral subarachnoid space is also warranted. In our series, this space was better identified on T2 SPACE images and was preserved in all cases of DAW and interrupted in all cases of SCH (Fig. 3). This is an intuitive but crucial finding given that a ventral cord herniation will, by definition, lead to interruption of the CSF plane. Although it is conceivable that a DAW may produce such severe mass effect to interrupt this CSF plane, we did not observe this in any of our patients. It is important to note that evaluation of the ventral subarachnoid space also may be limited by the slice thickness and resolution of standard MRI sequences. We suggest that high-resolution MRI be first performed to delineate the ventral subarachnoid space and a CT myelogram be used as the next step in the eventuality of the MRI being inconclusive. Our case series illustrates the value of high-resolution imaging via either T2 SPACE images or CT myelography in those instances in which conventional MRI is equivocal.

**Study Limitations**

The retrospective nature of our study and the small sample size are important limitations. The full gamut of standard and high-resolution MRI sequences was not available for all our patients, which limited a direct comparison of the accuracies of MRI and CT myelography. Also, no CSF flow-sensitive sequences were obtained in any of our patients. These may have some value in identifying the site of CSF flow interruption. For example, Chang et al. reported the use of cardiac-gated, phase-contrast, cine-mode MRI to precisely localize webs in 2 patients.\textsuperscript{3} Brugieres et al.\textsuperscript{1} employed phase-contrast MRI to demonstrate the absence of flow ventrally with preserved flow dorsally in 2 patients with SCH. We also did not include arachnoid cysts in our analysis. Arachnoid cysts are recognized by their marginated walls, smooth wide scalloping of the dorsal cord surface, and their tendency to present as slowly filling defects compared with the remainder of the subarachnoid space on myelography.\textsuperscript{12}

**Conclusions**

In summary, DAW and SCH both present as focal deformities of the thoracic spinal cord on imaging, but their differentiation has important surgical implications. In our small series, imaging reveals that the scalpel-shaped dorsal indentation and preservation of the ventral CSF plane are strongly suggestive of DAW while the C-shaped deformity and interruption of the ventral CSF appear to be typical of SCH. Although conventional MRI may be adequate in the characterization of these findings, high-resolution T2-weighted sequences and CT myelography are useful problem-solving tools.

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**References**


**Disclosures**

Dr. Sansur has served as a consultant to Medtronic, Globus, Stryker, and DePuy Synthes.

**Author Contributions**

Conception and design: Raghavan. Acquisition of data: Raghavan, Schultz, Steven, Gandhi. Analysis and interpretation of data: Raghavan, Schultz, Steven, Fischbein. Drafting the article: Raghavan, Schultz, Steven, Wessell, Fischbein, Sansur, Ibrahim. Critically revising the article: all authors. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: Raghavan. Study supervision: Raghavan.

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