INTRODUCTION

Spinal dural arteriovenous fistula (SDAVF) is the most common spinal vascular malformation, however it is still rare and underdiagnosed. Magnetic resonance imaging findings such as spinal cord edema and dilated and tortuous perimedullary veins play a pivotal role in the confirmation of the diagnosis. However, spinal angiography remains the gold standard in the diagnosis of SDAVF. Classic angiographic findings of SDAVF are early filling of radicular veins, delayed venous return, and an extensive network of dilated perimedullary venous plexus. A series of angiograms of SDAVF at different locations along the spinal column, and mimics of serpentine perimedullary venous plexus on MR images, are demonstrated. Thorough knowledge of SDAVF aids correct diagnosis and prevents irreversible complications.

Index terms: Central nervous system vascular malformations; Spinal cord diseases; Spine

Spinal Dural Arteriovenous Fistula: Imaging Features and Its Mimics

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The clinical course of SDAVF is slow and progressive (1). Initially, the symptoms can be nonspecific such as tingling pain, weakness, gait disturbance, paresthesia, and diffuse or patchy sensory loss (2-4). Back pain without specific radicular distribution commonly occurs with a burning sensation or cramping (3). These neurological deficits progress and become worse with time (5). Acute deterioration sometimes presents after exercise, prolonged standing, changing position, and even singing and eating serious morbidity and cause irreversible handicap.

Due to nonspecific symptoms, the neuroradiologist is often the first clinician to make the initial diagnosis based on MR imaging, with differential diagnoses causing confusion. In addition, spinal angiography is the gold standard and should be taken into consideration where SDAVF is highly suspected. A neuroradiologist should interpret the spinal angiography carefully in order to differentiate between SDAVF and other possible mimics.

Clinical Presentation

The clinical course of SDAVF is slow and progressive (1). Initially, the symptoms can be nonspecific such as tingling pain, weakness, gait disturbance, paresthesia, and diffuse or patchy sensory loss (2-4). Back pain without specific radicular distribution commonly occurs with a burning sensation or cramping (3). These neurological deficits progress and become worse with time (5). Acute deterioration sometimes presents after exercise, prolonged standing, changing position, and even singing and eating serious morbidity and cause irreversible handicap.

Due to nonspecific symptoms, the neuroradiologist is often the first clinician to make the initial diagnosis based on MR imaging, with differential diagnoses causing confusion. In addition, spinal angiography is the gold standard and should be taken into consideration where SDAVF is highly suspected. A neuroradiologist should interpret the spinal angiography carefully in order to differentiate between SDAVF and other possible mimics.
myelopathy. This may explain why the symptoms sometimes deteriorate after exercise or physical activities (6). The perimedullary venous plexus receives blood from radial veins draining the intramedullary circulation from the spinal cord dorsally in 80–90% of cases (21). Accumulated venous pressure can transmit through the perimedullary venous plexus in the caudo-cranial direction along the spinal cord (5). Extension can also go ventral and upwards to the cervical region or intracranial dural sinuses (22).

Radicular veins in the lower thoracic and lumbar region are fewer (23) and smaller in caliber, while venous drainage in the cervical region is divergent rather than convergent (16, 24). Therefore the lower thoracic and lumbar spinal cord is more vulnerable to hemodynamic changes, and the initial symptoms of SDAVF usually reflect dysfunction of the lower part of the spinal cord, that is, the conus medullaris (5, 25, 26). Cervical SDAVF is rare, and presents with hemorrhage more often than thoracolumbar counterparts, possibly due to a higher blood flow rate. SDAVF is often located at the upper cervical level with arterial feeding from the right-sided vessels, particularly the vertebral artery, and frequently in association with complex coincidental vascular lesions (8).

Tadié et al. (23), demonstrated a narrowing of the intervertebral veins at the point crossing the dura mater. It loses its vascular wall that is replaced by an arachnoid cuff by the dura mater, showing a zigzag fashion when exiting the dura. It was hypothesized that the zigzag fashion and

Epidemiology and Etiology

The exact etiology of SDAVF remains unclear. The majority of patients become symptomatic in middle age (15-18), suggesting that it is an acquired disease. Male predilection is also well known (18). Trauma is not considered a major causative factor in SDAVF due to the fact that trauma to the spine has been reported in merely 4% of patients (2). Furthermore, cervical SDAVF is rare despite the fact that the cervical spine is the most mobile segment of the spinal column and is more prone to trauma (2, 18).

As compared with spinal arteriovenous malformation (AVM), SDAVF is never located within the spinal parenchyma and seldom causes intramedullary hemorrhage. It occurs in the cervical region much less frequently, and patients with SDAVF are older than those with spinal AVM (15).

Pathophysiology

Typically, one (sometimes multiple) feeding artery from a radiculopial artery or a dural branch of a radiculopial artery enters an intervertebral or radicular vein and forms a SDAVF within the dorsal surface of the dural root sleeve in the intervertebral foramen (Fig. 1) (5, 15-17).

The feeding artery ‘arterializes’ the intervertebral vein, destroys the arteriovenous gradient and increases venous pressure, resulting in wall thickening and tortuosity of radial veins (also known as intramedullary veins), since intervertebral and radial veins share a common venous outflow (19, 20). Subsequently, chronic venous hypertension and stagnation decrease tissue perfusion and lead to edema of the spinal cord and progressive

Fig. 1. Illustration of SDAVF at lower thoracic level. Arteries are labeled with black background, while veins are labeled with white background. a = aorta, b = intercostal artery, c = dorsal branch of spinal artery, d = ventral branch of spinal artery, e = ventral epidural arcade, f = dural branch of ventral spinal artery, g = radicular artery, h = radiculomedullary artery, i = radiculopial artery, j = anterior spinal artery, k = dural branches of radiculopial artery, l = radial vein, m = perimedullary venous plexus, n = posterior radicular vein, o = anterior radicular vein, p = intervertebral vein, q = epidural venous plexus.
the narrowing of the intervertebral vein prevent blood flow going backwards to the intradural space, only permitting blood flow in a physiological direction. According to this observation, the “protective anti-flow back system” may be damaged in cases of SDAVF by congenital or acquired factors. However, patients with SDAVF may be asymptomatic despite the damaged “anti-flow back system” due to alternate venous outflow into ascending perimedullary veins, which open to the external paravertebral venous plexus via the upper cervical region. In these cases, SDAVF results in dilated perimedullary venous plexus but no venous hypertension or spinal cord edema (27). The average age of patients with asymptomatic SDAVF is roughly 10 years younger than that of patients with symptomatic SDAVF. Spontaneous angiographic conversion of progressive thrombosis and venous outflow obstruction related to aging has been reported in intracranial dural arteriovenous fistulae (28). It is noteworthy that the spontaneous angiographic conversion related to aging may also induce clinical symptoms in patients with asymptomatic SDAVF (27).

Diagnosis

Myelography could be used as a screening examination prior to spinal angiography (29). Typical findings include the presence of prominent vessels extending over an average of eight levels, as well as beading of the cauda equine (30). Computed tomography angiography is also considered as an efficient method of localization with its high resolution and high contrast, reducing the time required for angiography (31). However, magnetic resonance imaging (MRI) and catheter angiography techniques are important in the confirmation of the diagnosis when there is a clinical suspicion of a progressive myelopathy.

The cord edema is most clearly shown as hyperintensities with peripheral sparing on T2-weighted images (Fig. 2A) (30, 32-35). It could extend over an average of 5–7 vertebrae, with conus involvement in over 80% of cases, which are usually homogeneous (30, 34). A serpentine and dilated intervertebral vein or engorged perimedullary venous plexus as flow voids are another striking characteristic on T2-weighted images (Fig. 2A). Nevertheless, if the arterialized veins are too small to visualize, contrast enhancement may play a pivotal role at diagnosis (Fig. 2B, C). MR angiography can be helpful in identifying the possible arterial feeders, and avoid unnecessary superselective injections of all possible arterial feeders in more than two-thirds of patients (32, 35). With a negative MRI scan and no findings

Fig. 2. Characteristic diagnostic clues of SDAVF on MRI.
A. On sagittal T2-weighted spinal MR image, spinal cord hyperintensity (hollow arrows), and serpentine and dilated perimedullary venous plexus (white arrows) as flow voids are striking characteristics. B. On sagittal T1-weighted spinal MR image, tortuous and dilated perimedullary venous plexus is barely recognized (white arrows). C. Sagittal T1-weighted MR image with gadolinium enhancement clearly reveals tortuous and dilated perimedullary venous plexus (white arrows). SDAVF = spinal dural arteriovenous fistula
Fig. 3. Intracranial dural arteriovenous fistula with arterial feeding from meningeal branch of left vertebral artery of 47-year-old man with progressive quadriparesis associated with urinary and fecal incontinence and impotency.
A. Sagittal T1-weighted spinal MR image with gadolinium enhancement shows serpentine enhancement at anterior and posterior surfaces of cervical spinal cord and brainstem, as well as at tonsils (arrowheads).
B. Lateral view of left vertebral arteriogram demonstrates engorged inferior vermian vein (white arrows), which connects to veins at lower brainstem to drain into anterior and posterior spinal veins (hallow arrows). Reproduced with permission from Chen et al. Neuroradiology 1998;40:393-397 (64).

Fig. 4. Cervical SDAVF with feeding artery from right vertebral artery of 84-year-old man as incidental finding.
A. Right vertebral angiogram in frontal view at early phase demonstrates suspicious SDAVF (white arrows) with feeding artery directly from right vertebral artery (hallow arrow).
B. Dilated and tortuous draining vein is prominent at late phase (black arrows). SDAVF = spinal dural arteriovenous fistula
resembling SDAVF on MR angiography, SDAVF is unlikely to be the culprit of the symptoms (30, 32, 36).

Spinal catheter angiography, the gold standard in the diagnosis of SDAVF (18), should be considered if SDAVF is highly suspected due to its superior sensitivity compared with MR angiography. Ultimately, it is a treatable disease and severe morbidity can be avoided. The most frequent locations are the lower portion of the thoracic spine and the upper portion of the lumbar spine (2, 3), and 70% of SDAVFs are located on the left side in the Toronto series (4). The potential feeding artery can be as cranial as the intracranial vessels and as caudal as the lateral sacral artery or inferior hypogastric artery (Figs. 3-6) (37). Localization of these lesions may be difficult and challenging, especially in cases with cord edema distant from the arteriovenous shunt location.

During catheter angiography, the intercostal and lumbar arteries should be viewed as potential feeding arteries, in addition to the median and lateral sacral arteries and the deep cervical and ascending cervical arteries. Once an arterial feeder of SDAVF is localized, the tributaries at the ipsilateral and contralateral pedicle above and below the fistula should be catheterized to exclude the possibility of multiple feeding arteries to the fistula from the adjacent intercostal or lumbar arteries (29). Stasis of contrast material in the radiculomedullary arteries, especially the anterior spinal artery, can be found in catheter angiography followed by early filling of radicular veins and delayed

Fig. 5. Lumbar SDAVF with arterial feeding from left L2 lumbar artery of 53-year-old man presenting with bilateral weakness in lower extremities and urinary bladder frequency.
A. Sagittal T2-weighted spinal MR image reveals hyperintensity (hollow arrows) with flow voids (white arrows) mostly at posterior surface of spinal cord. B. Superselective angiography evidently shows feeding artery (hollow arrow), SDAVF (white arrow) and tortuous draining vein (black arrow). SDAVF = spinal dural arteriovenous fistula
venous return (5, 27). An extensive network of dilated perimedullary venous plexus is usually also visible.

**Mimics of SDAVF on MR Images**

Spinal cord edema and serpentine perimedullary venous plexus are two main characteristics of a SDAVF on MRI. In terms of spinal cord edema, the central hyperintensity on T2-weighted images resembles findings of anterior spinal artery infarction, myelitis, spinal cord neoplasms (38), or a persistent central canal if slit-like (39). In comparison, spinal cord edema due to a SDAVF is usually homogenous with a tapered margin and extends across an average of 5–7 vertebrae with conus involvement (34).

Differential diagnoses for serpentine perimedullary venous plexus include flow voids on T2-weighted images caused by severe spinal canal stenosis and spinal extradural arteriovenous fistulas (SEAVF), and artifacts caused by turbulent flow of cerebrospinal fluid (CSF).

Severe spinal canal stenosis may display an engorged and tortuous vein, which is usually across only 1–2 vertebrae, along with a tangle of redundant nerves and vessels just proximal to the level of stenosis on myelography or MR myelography (Fig. 7). MRI helps to avoid confusion and shows no spinal cord edema or serpentine flow voids on the dorsal surface of the spinal cord.

Dural or epidural arterial branches drain primarily into epidural venous plexuses and form a SEAVF that is even rarer than SDAVF. To recognize and differentiate a SEAVF from a SDAVF is essential due to the fact that the ventral location to the dural sac and the arterialized venous lakes are challenging in hemostasis during surgery. The arterial feeders of SEAVF could originate from multiple levels, and the engorged epidural venous plexuses may cross the midline and fill multiple compartments, showing characteristic diamond-shaped plexuses (Fig. 8). A SDAVF, however, usually has a single feeding artery and drains intradural veins without filling the epidural space.

Turbulent flow of CSF moving from the ventral to the dorsal part of the subarachnoid space, combined with cardiac and respiratory related pulsatile CSF flow, leads to signal loss seen clearly on sagittal T2-weighted images, which is a frequently encountered artifact (40). It mostly occurs in children due to a relatively wider CSF space at the thoracic level, as well as patients with residual dorsal arachnoid septum posticum or spinal arachnoid cysts.

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**Fig. 5. Lumbar SDAVF with arterial feeding from left L2 lumbar artery of 53-year-old man presenting with bilateral weakness in lower extremities and urinary bladder frequency.**

C. Angiogram of left L2 lumbar artery in frontal view before treatment demonstrates fistula (white arrow) and tortuous vein (hollow arrow). D. Angiogram of left L2 lumbar artery in frontal view after treatment reveals result of successful endovascular embolization without opacification of fistula or tortuous vein.
The signal loss is bulky, discontinuous, and prone to be observed at the thoracic level (Fig. 9).

It is intriguing to find prominent spinal arteries mimicking dilated perimedullary venous plexus on MR images. T2-weighted sagittal MR images show no spinal cord edema. Spinal angiography demonstrates the anatomy precisely, and proves the diagnosis of prominent spinal arteries (Figs. 10, 11).

**Treatment**

Spontaneous closure of a SDAVF without treatment is extremely rare (41). Treatment is essential because an untreated SDAVF can progress to serious morbidity and cause irreversible handicap.

Treatment options of a SDAVF include endovascular embolization and surgical ligation of the fistula. Endovascular embolization is less invasive. First of all, superselective catheterization of the feeding radiculopial artery should be carried out. The goal of treatment of SDAVF is to occlude the nidus as well as the first 1–2 cm of the proximal draining vein of the fistula (42). Embolization of only the distal artery may relieve the symptoms temporarily but almost always inevitably leads to recurrence due to numerous collaterals (43). Embolization with liquid polymers such as N-butyl 2-cyanoacrylate (NBCA), which is commonly referred to as glue, is recommended for a higher success rate.
rate of 44–100% (4, 44-46). On the contrary, particles such as polyvinyl alcohol are thought to be related to the high recurrence rate of 30–93% (47) due to early recanalization, and coils and gelfoam are contraindicated for the same reason (5). The concentration of NBCA is a significant predictor of successful endovascular embolization. A higher concentration of NBCA presumably polymerizes faster and does not penetrate well. A lower concentration of NBCA polymerizes slower, but may not penetrate well either due to the higher viscosity resulting from the higher amount of iophendylate (lipiodol). The optimal concentration of NBCA is reported to be approximately 20–30% in order to minimize the obstacles of fast polymerization and high viscosity (48). In our experience, it is best accomplished with a mixture that has a long polymerization time such as NBCA and lipiodol in the ratio 1:4 (20%), which may be adjusted to 1:3 (25%) if the blood flow is fast. The mixture can be injected in a consecutive column or with a so-called sandwich or push technique. The direction of flow during final microcatheter injection is another determining factor in procedural success. When the flow is predominantly towards the fistula, the glue mixture has a higher chance of reaching the draining vein (48).

Ethylene vinyl alcohol (Onyx) is a liquid embolic material
Fig. 10. Prominent anterior spinal artery mimics SDAVF.
A. Sagittal gadolinium-enhanced MR image demonstrates prominent vessels on anterior surface of spinal cord (white arrows). B, C. Lateral (B) and frontal (C) views of left T12 intercostal arteriogram reveal prominent artery in middle of anterior surface of spinal cord (black arrows) with characteristic hairpin turn, which is anterior spinal artery supplying from artery of Adamkiewicz. No SDAVF was found based on this study. SDAVF = spinal dural arteriovenous fistula.

Fig. 11. Prominent left posterior spinal artery mimics SDAVF.
A. Left L1 lumbar arteriogram shows longitudinal artery without hairpin turn. B, C. On coronal (B) and sagittal (C) three-dimensional CT angiograms, artery is paramedian and dorsal to spinal cord (hollow arrow), proven to be prominent left posterior spinal artery. White dotted line demonstrates midline. SDAVF = spinal dural arteriovenous fistula.
dissolved in dimethyl sulfoxide that polymerizes soon after contact with the bloodstream (49). Advantages of Onyx include lower viscosity, which facilitates venous penetration and decreases the risk of microcatheter retention after embolization, and theoretical permanent polymerization with empirical evidence of a low rate of recanalization of onyx-embolized aneurysms at the 5-year follow-up (50). Disadvantages of Onyx include the requirement for compatible catheters, higher expense, and the relative lack of availability in many centers (42). Transarterial Onyx embolization has been accepted as a promising therapeutic option for impressive feasibility and safety over NBCA, while long-term results remain controversial (42, 51, 52).

Endovascular embolization is not always viable. If a spinal cord-supplying artery arises from the same pedicle as the feeder to the shunt, such as the great anterior radiculomedullary artery that is also known as the artery of Adamkiewicz, embolization may cause spinal cord ischemia (Fig. 12). Furthermore, technical barriers such as severe atherosclerosis, arterial dissection of the feeding arteries, or extensive collateral vessels may halt superselection of target vessels.

Surgical treatment should be considered if endovascular embolization fails. Currently, ligation of the draining vein is recommended (53), and has been proven to be as effective as total removal of the draining vein (54).

The success rate of surgery (98%) is higher than that of embolization (46%) (55). However, embolization is less invasive and more efficient when only a single arterial feeder is shown, and can be used preoperatively to label the feeding artery (5). Generally, the prognosis depends on the duration of symptoms, pretreatment disabilities, and degrees of obliteration of the fistula and draining veins, however, no reliable prognostic factors have yet been identified (45, 56-58).

It has been reported that 55% of patients had improved gait after microsurgery, 11% deteriorated, and the rest remained stationary, with only 33% having improvement in micturition. These results are compatible with vascular anatomy and the distribution of motor and sensory pathways in the spine. Other studies have also yielded a similar conclusion that patients benefit most from the treatment, generally endovascular embolization, in motor and pain symptoms, followed by sensory disturbance, and sphincter and erectile dysfunctions (56, 58-61). It is reasonable to presume that endovascular embolization could be as effective in patients with complete obliteration of the fistula as in those who undergo surgery.

If there is no improvement within 4–6 weeks following treatment, or even clinical deterioration during follow-ups, repeat MRI should be performed. Once MRI distinctly demonstrates persistent flow voids along the spinal cord in a patient with a poor response to the previous treatment, repeat angiography is indicated (62). Residual fistula caused by inadequate occlusion or recanalization, and rarely but not impossibly a second fistula, should be looked for carefully. Therefore, follow-up angiography is recommended...
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to be carried out 2–3 months after surgical or endovascular treatment, due to the fact that clinical improvement starts to reach a plateau around that time. High suspicion of residual fistula prompts an earlier follow-up angiography (63).

CONCLUSION

Spinal dural arteriovenous fistula is rare but frequently underdiagnosed since the clinical presentations resemble other more common causes of myelopathy. The disease is treatable but can progress to serious morbidity and irreversible damage. That is, the earlier the diagnosis, the better the outcome. MR imaging and spinal angiography should be considered under high clinical suspicion. Subsequently, prompt endovascular embolization or surgical ligation should be performed according to the conditions of the patient and characteristics of the fistula. Thorough knowledge regarding SDAVF may enable clinicians and neuroradiologists to diagnose the disease in the early stages.

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