MR Diffusion Tensor Imaging and Fiber Tracking in Spinal Cord Arteriovenous Malformations: A Preliminary Study

A. Ozanne, T. Krings, D. Facon, P. Fillard, J.L. Dumas, H. Alvarez, D. Ducreux and P. Lasjaunias

AJNR Am J Neuroradiol 2007, 28 (7) 1271-1279
doi: https://doi.org/10.3174/ajnr.A0541
http://www.ajnr.org/content/28/7/1271
MR Diffusion Tensor Imaging and Fiber Tracking in Spinal Cord Arteriovenous Malformations: A Preliminary Study

BACKGROUND AND PURPOSE: Diffusion tensor imaging (DTI) of the spinal cord in patients harboring spinal arteriovenous malformations (AVMs) was carried out to evaluate the feasibility of this new technique to determine the displacement of the spinal cord tracts and to correlate morphologic and functional DTI data (fractional anisotropy [FA] and apparent diffusion coefficient [ADC]) with the clinical symptoms.

MATERIALS AND METHODS: Nine patients with spinal cord AVMs were investigated at 1.5T using a sagittal spin-echo single-shot echo-planar generalized autocalibrating partially parallel acquisition diffusion-weighted imaging sequence. ADC and FA maps were computed in different regions of interest (both above and below the nidus), and tractography was used to visualize the course of the tracts. The data were correlated with the clinical symptoms and compared with 12 healthy control subjects.

RESULTS: At the level of the nidus, tracts were normal, shifted, separated, or interrupted but not intermingled with the nidus. Interruption of the tracts was coherent with the clinical symptoms. In patients with severe neurologic deficits, FA values caudal to the nidus showed a reduced anisotropy consistent with loss of white matter tracts.

CONCLUSIONS: We demonstrate that AVMs may interrupt, displace, or separate the fiber tracts and that clinical symptoms may be reflected by the quantitative FA results and the morphologic loss of fibers distant to the lesion. DTI with fiber tracking offers a novel approach to image spinal cord AVMs and may open a window to understand the complex pathophysiology of these lesions.

In recent years, many studies concerning MR diffusion tensor imaging (DTI) and fiber tracking have been published concerning both the normal aspect and various pathologies in the brain, including brain arteriovenous malformations (AVMs). Concerning the spinal cord, this MR technique is more difficult to apply because of the following: 1) the small volume of the spinal cord compared with the brain, 2) difficulties in obtaining a homogeneous magnetic field inside the spinal canal, and 3) susceptibility artifacts related to the echo-planar imaging (EPI) technique used typically. Therefore, there is paucity of data concerning spinal cord DTI. Studies reported thus far describe DTI findings in compressive disease, chronic myelopathy, and multiple sclerosis. However, DTI has not yet been reported in vascular malformations affecting the spinal cord. AVMs of the spinal cord may give rise to an increase of the extracellular water because of venous congestion and/or a loss of white matter fibers (eg, because of Wallerian degeneration after hemorrhage) in the normal-appearing cord distant to the nidus. To further differentiate the above-mentioned possible effects of spinal cord AVMs on the white matter, it appears appealing to use DTI because its principle is based on the anisotropic diffusion of extracellular water molecules within the anisotropic environment of white matter fibers.

One purpose of this study was to analyze the quantitative results of the apparent diffusion coefficient (ADC) and the fractional anisotropy (FA) that were measured both cranial and caudal to the nidus to evaluate whether these data can give information on the complex pathophysiology of spinal cord AVMs. Apart from hemorrhage, spinal vascular malformations can lead to neurologic deficits via various pathologic mechanisms, including venous congestion and mass effect. For a particular AVM, however, it is not known which of these pathologic mechanisms is the most relevant one, though this information might affect the therapeutic approach (ie, partial targeted endovascular treatment for compartments likely to rehemorrhage, endovascular flow reduction to treat venous congestion, or reduction of size of venous pouches to treat the mass effect). We hypothesize that the quantitative data could shed light on the pathologic mechanism to be encountered in a particular AVM, because, for example, mass effect may lead to an increase in anisotropy (because of compression of fibers), whereas cord edema because of venous congestion may lead to a decrease (because of separation of fibers). The main goal of the study, however, was to evaluate the feasibility of this new technique to determine the displacement of the spinal cord tracts and to correlate morphologic and functional DTI data (FA and ADC) with the clinical symptoms.

Materials and Methods

Subjects
Twelve patients with spinal cord AVM referred to our institution for endovascular treatment by transarterial embolization between March 2003 and December 2004 underwent MR imaging before treatment. Three patients were excluded from the present series: 1 because of
previous surgery of the spinal cord and 2 because of excessive movement artifacts during scanning. The remaining 9 patients (5 male and 4 female patients) had a mean age of 23.7 years (range: 13–57 years). Clinical and demographic data are described in Table 1. The mean time elapsed between the onset of symptoms and MR examination was 25.7 months, ranging from 3 months to 8 years. Three patients were referred to us less than 6 months after the onset of symptoms. Six patients had an acute onset of symptoms (<48 hours), and 3 presented with chronic progressive symptoms. In 1 patient (patient 3), 2 previous embolization sessions had been performed 4 and 3 years before DTI imaging, respectively; in the other patients, no previous embolization had been performed.

The severity of the symptoms was evaluated with the clinical functional score that was described previously by McCormick et al. In this grading system, grade I is designated to patients whose neurologic examination is normal or who demonstrate a mild deficit without impairment of the function of any limb. In patients at grade II, sensorimotor deficits of limbs and/or severe pain and/or sensory deficits responsible for an impairment of the quality of life but compatible with independent and self-sufficient walking are present. Grade III is characterized by severe sensorimotor deficits of the inferior limbs or disabling sensorimotor deficits of the superior limbs. In grade IV, patients have deficits that render them totally dependent. Three patients had none or a mild deficit (grade I), 4 patients had a deficit resulting in a decrease in the quality of life (grade II), and 2 patients had a deficit resulting in total dependence in activities of daily living (grade IV). These latter 2 patients (patients 2 and 7) were found to have had an hemorrhagic event, as assessed by the combination of clinical and imaging data. As our routine imaging protocol before embolization included MR imaging, we did not deem it necessary to obtain ethics committee approval to add the DTI sequence; however, informed consent by all of the patients and institutional review board approval for chart and record review were obtained.

**MR Imaging Technique**

Imaging was performed on a 1.5T MR imaging system with actively shielded magnetic field gradients (maximum gradient: 40 mT/m). The protocol began with conventional sequences: acquisition of a T2-weighted coronal scout followed by a sagittal T2-weighted fast spin-echo (SE) sequence (FOV: 30 × 30 cm; image matrix: 640 × 384; section thickness: 3 mm; TR/TE: 4370 ms/119 ms), a sagittal T1-weighted SE sequence (FOV: 30 × 30 cm; image matrix: 576 × 346; section thickness: 3 mm; TR/TE: 400 ms/12 ms), and an axial T2-weighted gradient-echo sequence (FOV: 20 × 15 cm; image matrix: 256 × 134; section thickness: 4 mm; TR/TE: 1330 ms/27 ms).

DTI was performed using a sagittal SE single-shot echo-planar generalized autocalibrating partially parallel acquisition (GRAPPA) diffusion-weighted imaging sequence with a 2-fold acceleration factor; b values (b = 0 and 500 s/mm²) along 6 noncollinear gradient directions (DTIb500) for patients 2, 4, 6, 7, 8, and 9 and along 25 directions (DTIb2500) for the remaining patients (1, 3, and 5) were applied18,19 (FOV: 17.9 × 17.9 cm; image matrix: 128 × 128; 12 sections with section thickness of 3 mm; nominal voxel size: 1.4 mm × 1.4 mm × 3 mm; TR/TE: 4600 ms/73 ms for both DTIb2500 and DTIb500; the acquisition number was 6). Patients were asked to avoid swallowing and moving the head and limbs during diffusion-weighted imaging.

**Image Analysis**

Image analysis was performed on a voxel-by-voxel basis by using dedicated software (DPTools, http://www.fmritools.org). In addition to

---

**Table 1: Clinical data**

| Patients | Age (Years) | Sex | Presenting Symptoms | Time Elapsed between Onset of Symptom and Imaging | Clinical Status before Imaging | McCormick Score | Level of the Nidus | Hyperintensity Hemorrhage | T2 Hemorrhage | DTI25d | DTI6d |
|----------|-------------|-----|---------------------|---------------------------------------------|--------------------------------|-----------------|-------------------|--------------------------|----------------|--------|-------|
| 1        | 57          | F   | Progressive paresthesia followed by motor weakness of lower extremities | 18 months | Normal | C2-C3 | C3-C7 | + | + | | |
| 2        | 29          | F   | Paraparesis developed within 48 hours | 4 months | Paraparesis of lower limbs | C2-C3 | | | | + | |
| 3        | 23          | M   | Paresthesia of superior limbs | 3 months | Paresthesia of upper limbs | C2-C3 | | | | | |
| 4        | 19          | M   | Paraparesis developed within 48 hours | 6 months | Paraparesis of all extremities | Normal | | | | | |
| 5        | 14          | M   | Transitory paraplegia | 4 months | Paraparesis of inferior limbs | C1-C2 | | | | | |
| 6        | 21          | F   | Acute tetraplegia | 8 years | Paraparesis of inferior limbs | C2-C3 | | | | | |
| 7        | 18          | M   | Paraplegia developed within 24 hours | 6 months | Paraplegia and sphincterian deficit | C2-C3 | | | | | |
| 8        | 35          | F   | Acute hemiparesis | 2 years | Hemiparesis | C2-C3 | | | | | |
| 9        | 13          | M   | Acute hemiparesis | 2 years | Hemiparesis | C2-C3 | | | | | |

Note: —F indicates female; M, male. McCormick clinical functional score: grade I: neurologic examination normal or mild deficit without impairment of the function of any limb and normal walking; grade II: sensitivomotor deficit responsible of an impairment of the quality of life but compatible with independent and self-sufficient walking; grade III: severe sensitivomotor deficit of any limb and incapable to walk; and grade IV: deficit of inferior and/or superior limbs responsible for a total dependence.
the 2D parametric FA color maps, 3D white matter fiber tracts were created using the principal diffusion directions method after oversampling and interpolation of the DTI dataset (n = 1 for 6 directions; n = 10 for 25 directions). In the principal diffusion directions method, the eigenvector corresponding with the largest eigenvalue is extracted from the diffusion tensor field generated from the DTI datasets in the region where the diffusion is linear, typically for a FA thresholding value of 0.18 and an angulation threshold of 45° (to prevent fibers from sudden transition and to keep tracking based on the connectivity of the neighborhood), as described previously.

When morphologically analyzing the fiber tracking (FT), we qualitatively described the tracts as normal, as defined previously, shifted, separated, or interrupted (partially, i.e., across parts of the axial plane, or completely, i.e., across the entire axial plane). Concerning quantitative data, we made measurements on the FA and ADC maps. We placed regions of interest (ROIs) centrally within the cord to avoid partial volume effects with the CSF using a technique described previously (Fig 5). As specified in Table 2, we tried to locate the ROIs for each patient. In 2 patients, the AVM nidus was located at the cervicomedullary junction; therefore, measurements cranial to the nidus were not feasible;
in 1 patient, measurements caudal to the nidus were not possible, because the FOV was too small. Because in some subjects, T2 abnormalities were present, whereas in others, the cord appeared normal, we placed additional ROIs in the area of T2 hyperintensity and in the normal-appearing cord (Table 2). Measurements at the level of the nidus were only possible in a single subject, otherwise, quantitative measurements at the level of the nidus were not feasible because of susceptibility artifacts, thrombosis, glue artifacts, enlarged venous pouches, residua of old hemorrhages, or small volume of the residual cord. If the T2 hyperintensity had ill-defined margins with concomitant swelling of the cord, venous congestion was presumed to be the cause of the hyperintensity compared with gliosis or nonspace-occupying syringohydromyelia if the margins were well delineated and the volume of the cord was normal or decreased, a state that we will further refer to as “cavitation.”

Statistical Analysis

We used the method of Z-scores to compare the values of the patients with those of 11 healthy volunteers (8 men and 3 women; mean age: 36.7 years; range: 20 to 67 years) who were investigated with the same imaging protocol. Their FA and ADC values were 0.748 with a range of 0.7 to 0.8 for FA and 1.00 $10^{-3}$ with a range of 0.77 to 1.25 $10^{-3}$ for the ADC. FT showed the main white matter tracts (posterolateral corticospinal tracts and posterior lemniscal tracts) in all of the volunteers. The values of the AVM were considered abnormal when not included in the 95% confidence interval [Z-score $> 2$, $Z = (\bar{x} - \mu)/\sigma$], with $x =$ value, $\mu =$ mean, and $\sigma =$ SD of the healthy volunteers.

Results

The most illustrative cases are reported in Figs 1-6 and show mainly the morphologic aspect of the fiber tracts compared with conventional MR imaging and digital subtraction angiography findings. In the following, we detail both the FT and the quantitative data of DTI in comparison with morphologic and clinical data.

DTI Results in Relation to the MR Morphology

At the level of the nidus, the fibers were separated, shifted, interrupted, or normal compared with the morphology of fibers in healthy control subjects. No fibers running through the nidus itself could be detected, comparing the results of the FT data with the conventional imaging data.

Distant to the nidus, congestive edema was present in 1 patient: FT showed separated fascicles, with global enlargement of the beam of tracts. There was a slight rarefaction of tracts (Fig 1). We found significantly decreased FA (0.42/$Z = -11.83$) and normal ADC values (0.955 $10^{-3}$/Z = -0.34).

Cranial or caudal to the nidus, a cavitation as explained above was present in 3 patients (Figs 2 and 5). FT demonstrated a loss of tracts with global thinning of the beam. The remaining tracts were still grouped into fascicles and were separated from the midline. A significantly decreased mean FA of 0.42/$Z = -11.7$ (FA range: 0.36–0.50) and a normal mean ADC of 1.13 $10^{-3}$/Z = 1.02 (ADC range: 0.79–1.45 $10^{-3}$) were found.

In the segments of the cord distant to the nidus without T2 hyperintensity where draining veins were present, FA values were marginally decreased compared with those segments where no draining veins were visualized. DTI in normal-appearing cord above and below the nidus and distant to the T2 hyperintensity also showed slightly lower FA values in the group of patients compared with the normal cord of healthy volunteers (mean FA: 0.64 $Z = -3.7$), whereas the ADC was normal without statistically significant differences between both groups.

DTI Results in Relation to the Clinical Symptoms

In addition to the above-mentioned correlations of FA and ADC results to MR imaging data, we also performed a correlation with the clinical symptoms. At the level of the nidus, 1 patient with a severe neurologic deficit (Mc Cormick IV) who became symptomatic because of a spinal hemorrhage had a complete interruption of the tracts (Fig 6). Five patients had partially interrupted tracts (Figs 3 and 5): 4 of
them had a score of McCormick I or II without any obvious previous hemorrhage; the fifth had a more severe clinical score (McCormick IV) and a previous hemorrhage. In 2 patients, the tracts were separated but not interrupted at the level of the nidus, and the clinical score was good (McCormick I and II, respectively). In 1 patient, the tracts were shifted without focal interruption, and the clinical status was normal (Fig 4).

Under the assumption that FA and ADC in the normal-appearing cord caudal to the nidus may be altered because of the AVM, we investigated quantitative DTI results caudal to the cord. Four patients had a good clinical score (McCormick I). In these patients, the FA values were normal compared with the group of healthy control subjects (mean FA: 0.74/0.51), whereas the ADC values were slightly decreased (mean ADC: 0.849 10^-3/1.0). Patients with neurologic symptoms had decreased FA values, both compared with the group of healthy volunteers and with the group of patients with good clinical scores. Three patients with a score of II had a mean FA of 0.68 (Z = −2.5),
whereas ADC values were similar compared with patients with a score of I (mean ADC: 0.861 \times 10^{-3}; Z = -1.0). In the 2 patients with severe neurologic impairment (McCormick IV), FA values caudal to the AVM could be measured for technical reasons in only 1 patient. In this patient, the decrease in FA values found caudal to the nidus in the normal-appearing cord was most pronounced with a mean FA of 0.55 (Z = -6.8), whereas ADC values were normal (mean ADC: 1.014 \times 10^{-3}; Z = -0.11).

Quantitative DTI results in the normal-appearing cord were also investigated with regard to the duration of symptoms. However, here we found no difference between the group of patients who were imaged with a longer latency and the patients imaged in close temporal relation to their symptom onset.
Discussion

Background

Spinal cord AVMs are usually diagnosed by conventional MR imaging showing an intramedullary nidus and dilated perimedullary draining veins.\textsuperscript{9,23,24} MR imaging can also depict potential complications of AVMs, such as hemorrhage (hematomyelia or subarachnoid hemorrhage) or venous congestion because of altered venous drainage. This congestive edema of the spinal cord may extend over multiple segments and is due to an increase of interstitial water. Chronic venous congestion leads to slowly progressive neurologic symptoms and spinal cord atrophy. In patients with hematomyelia, AVMs may lead to variable neurologic deficits resulting from direct damage to the gray and white matter.\textsuperscript{10} Endovascular treatment by transarterial embolization with glue is aimed first at excluding those parts of the nidus that harbor identifiable angioarchitectural risk factors, such as intranidal false aneurysms or drainage into venous ectasias or stenosis. A second goal is to reduce the flow in the nidus and subsequently in the draining veins to decrease the venous congestion.\textsuperscript{25} MR imaging performed after embolization may, therefore, show complete regression of the T2 hyperintensity. Our series of 9 patients with spinal cord AVM investigated with DTI presented with various clinical and imaging findings in terms of the presence of hemorrhage, venous congestion, and the time elapsed between the onset of the symptoms and the admission. Although this group of patients allows us to evaluate the use of DTI in a wide variety of different pathologic mechanisms present in AVM, it makes the investigated group of patients rather heterogeneous. Still, there were certain characteristic findings that warrant further discussion.

$\text{DTI at the Level of the Nidus}$

At the level of the nidus, FT may give additional information compared with the conventional morphologic MR images. As could be demonstrated in our series, the morphologic aspects of the fiber tracts could be demonstrated in all 9 of the patients, though the exact delineation among the nidus, the intrinsic draining veins, and the white matter was difficult to perceive because of low spatial resolution. In most patients, FT did not show focal interruption of tracts at the level of the nidus. This is coherent with the notion that an AVM, being in the subarachnoid or subpial space, does not “infiltrate” the normal spinal cord but in fact displaces or separates the fiber tracts. For brain AVMs, this has been recently illustrated by Maruyama et al,\textsuperscript{26} who, by using FT, were able to show that corticospinal tracts in patients with rolandic AVMs are separated or displaced, but there are no tracts running through the vascular malformation. In our series, continuous tracts were correlated with a good clinical status. If interruption of tracts was present, clinical symptoms corresponded with the interrupted tract (Fig 3: patient with loss of sensation and interrupted tracts at the T2-level, arrow). For brain AVMs, this has been recently illustrated by Maruyama et al,\textsuperscript{26} who, by using FT, were able to show that corticospinal tracts in patients with rolandic AVMs are separated or displaced, but there are no tracts running through the vascular malformation. In our series, continuous tracts were correlated with a good clinical status. If interruption of tracts was present, clinical symptoms corresponded with the interrupted tract (Fig 3: patient with loss of sensation and interrupted posterior white matter fibers). The cause of the interruption of the tracts may be because of one of the following: 1) the real loss of white matter fibers; 2) magnetic susceptibility artifacts because of the hemoglobin degradation products, the nidus, or the draining veins; or 3) an increase of free water in the vicinity of the nidus with subsequent decrease in anisotropy. When the nidus was not located inside the cord but close to its surface, the added value of the FT was not so evident, because T2 sequences could already demonstrate sufficiently the shifted cord distinct from the nidus. Still, FT was able to demonstrate the continuity of white matter bundles that were shifted by the nidus. For those patients who did not present with hematomyelia, there was no obvious discordance between the clinical data and the appearance of the tracts (preserved, shifted, or interrupted) at the level of the nidus as depicted by FT. Problems of FT arise, however, in patients with hemorrhage, because the interruption of tracts may be the consequence of susceptibility artifacts because of hemoglobin.
degradation products and may not be related to damage of the white matter from the hematomyelia. Furthermore, susceptibility artifacts from thrombosed pouches, pulsation, and flow artifacts render FT difficult to interpret. Evaluation of routine anatomic sequences in addition to FT images is, therefore, recommended.

**DTI in Venous Congestion**

In the single patient who presented with venous congestion, we were able to show that the tracts were heterogeneous separated and grouped into fascicles that may correspond with the anatomic fascicles. Two mechanisms may explain this aspect: a heterogeneous loss of white matter fibers predominating at the periphery of the white matter fascicles or a heterogeneous repartition of the increase of water in the interstitium related to the angioarchitecture of the intrinsic vascular network.

DTI is mainly based on the directed diffusion of extracellular water and is related to the interaction between the white matter fibers and the extracellular water. During venous congestion, the increase in extracellular water will, therefore, lead to a decrease of the FA while the fibers are still intact. FA is more sensitive than SE sequences to detect an increase of water, as Facon et al. had already reported in compressive pathology of the spinal cord. Depending on the intensity and the duration of the congestion, however, white matter fibers may become irreversibly damaged over time, leading to a further decrease of FA values. Because there has been only 1 patient in our series presenting with venous congestion distant to the nidus, our results have to be interpreted with caution. More patients, especially those with early or mild venous congestion, need to be studied to further assess the effect of venous congestion on FA.

**DTI in Cavitation**

In patients presenting with sharply delineated T2 hyperintensities of the central spinal cord representing either gliosis or non-space-occupying syringohydromyelia, FT showed a global thinning of the beam. We presume that this thinning is related to a true decrease of the white matter bundles in the vicinity of the lesion as shown by the concomitant decrease of the size of the cord on T2-weighted images. The decrease in FA at these levels further confirms the loss of fibers, whereas the normal mean ADC speaks in favor of a gliosis instead of a syringohydromyelia as the most probable morphologic substrate of the observed imaging findings. Nevertheless, quantitative data must be interpreted with caution because of limited spatial resolution of DTI with subsequent partial volume averaging within our longitudinal ROIs. The measurement with the ROI that we applied cannot distinguish between the different compartments of the spinal cord (ie, gray and white matter) depicted in high-resolution axial sections. Higher field strengths are necessary to overcome this problem, as shown in animal studies at 7T, where the FA values for normal gray and white matter are 0.59 and 0.92, respectively. Because we placed our ROIs in the sagittal plane, both white and gray matter areas are likely to be averaged in various and unknown proportions for different patients, thereby affecting the quantitative results considerably.

**DTI Findings Related to Clinical Aspects**

Caudal to the nidus within the normal-appearing cord, we were able to show a decrease of FA values in patients with poor clinical scores compared with both patients with good clinical scores and the group of healthy control subjects. These findings may be related to the loss of fibers because of wallerian degeneration, which has already been described for the brain after stroke. After cerebral infarction, a reduced anisotropy within the corticospinal tract remote from the lesion was found 2–6 months after ischemic stroke, highlighting the potential of DTI to detect and monitor the structural degeneration of fiber pathways.

A significant limitation of our conclusions is because of the low number of subjects and the heterogeneous patient population with a small number of patients with the respective pathologies (1 patient with an edema and 3 patients with what we call a cavitation). Therefore, we cannot decide whether the ADC values that we have seen in these patients truly reflect biological phenomena or whether they constitute an artifact that we cannot fully explain. More patients are needed to fully characterize the ADC values under these circumstances in the spinal cord. One might, for example, presume that, in venous congestion, not only an increase in water may be present but that a true cellular infiltration can occur. Because of the longstanding nature of the disease in some patients, one might also think of gliosis as the reason for the normal ADC values. In cases of cavitation, arachnoidal adhesions may be present that account for a non-elevated ADC. Potential sources of artifacts that may explain our findings are partial volume averaging, artifacts from the acquisition technique, and distortions because of local field inhomogeneities. However, our primary aim was to evaluate whether DTI of the spinal cord is feasible in patients harboring spinal vascular malformations. The complex and different pathophysiologies including venous congestion and hematomyelia and the different time intervals between symptom onset and imaging interfere with the interpretation of the results. Facing a loss of tracts, it remains impossible to differentiate a true loss of white matter fibers from the increase of interstitial-free water. The calculated fiber trajectories and ROI statistics therefore do not necessarily reflect true pathology. Susceptibility artifacts (because of hemoglobin degradation products or vascular structures) and the heterogeneity of the magnetic field in the spinal canal are additional problems. Although we were able to ameliorate our imaging sequences by changing the protocol from DTI_{4d} to DTI_{3d}; a single-shot EPI sequence still suffers from strong susceptibility distortions. Using parallel imaging techniques, this susceptibility distortion may be reduced; still, GRAPPA parallel imaging as used in this article may cause unfolding artifacts that may give rise to a pseudointerruption of spinal cord fibers. These problems may potentially be overcome with the use of new MR acquisition techniques, such as multishot SE DTI or a reversed gradient approach (that reduces in-plane voxel displacements by correcting distortions in the phase-encoding direction) or trying higher acceleration factors that may reduce some of these susceptibility and geometric distortion artifacts and that warrant further studies in this field.
Conclusions
In this short series we were able to show the following: 1) DTI of the spinal cord is possible in the presence of vascular malformations; 2) AVMs at the level of the nidus can interrupt, displace, or separate the fiber tracts, whereas, distant to the nidus, the effects of congestion or gliosis on the tracts can be perceived; 3) the clinical score can be reflected by the interruption of fibers close to the nidus and by a loss of fibers distant to the lesion; and 4) the quantitative results, especially the FA values, may be related to the clinical symptoms. We conclude that DTI with FT offers a novel approach to image spinal cord AVMs and opens a window to understand the complex pathophysiology of these lesions.

Acknowledgments
We thank one anonymous reviewer for his helpful and constructive critique. Moreover, we are deeply indebted to Soke M. Chng, MD, and Sean Cullen MD, for final proofreading and editing of the article.

References
1. Yamada K, Kizu O, Ito H, et al. Tractography for arteriovenous malformations near the sensorimotor cortices. AJNR Am J Neuroradiol 2005;26:598–602
2. Ries M, Jones RA, Doussset V, et al. Diffusion tensor MRI of the spinal cord. Magn Reson Med 2000;44:884–92
3. Clark CA, Werring DJ, Miller DH. Diffusion imaging of the spinal cord in vivo: estimation of the principal diffusivities and application to multiple sclerosis. Magn Reson Med 2000;43:133–38
4. Bammer R, Augustin M, Strasser-Fuchs S, et al. Magnetic resonance diffusion tensor imaging for characterizing diffuse and focal white matter abnormalities in multiple sclerosis. Magn Reson Med 2000;44:583–91
5. Cercignani M, Horsfield MA, Agosta F, et al. Sensitivity-encoded diffusion tensor MR imaging of the cervical cord. AJNR Am J Neuroradiol 2003;24:1254–56
6. Facon D, Oranne A, Fillard P, et al. MR diffusion tensor imaging and fiber tracking in spinal cord compression. AJNR Am J Neuroradiol 2003;26:1587–94
7. Demir A, Ries M, Mooren CT, et al. Diffusion-weighted MR imaging with apparent diffusion coefficient and apparent diffusion tensor maps in cervical spondylotic myelopathy. Radiology 2003;229:37–43
8. Valasinas P, Rocca MA, Agosta F, et al. Mean diffusivity and fractional anisotropy histogram analysis of the cervical cord in MS patients. Neuroimage 2005;26:822–28
9. Berenstein AL, Lasjaunias P, ter Brugge, KG. Spinal arteriovenous malformation. In: Surgical Neuroangiography. 2nd ed. Berlin: Springer-Verlag; 2004:37–843
10. Kringis T, Mull M, Gilsbach JM, et al. Spinal vascular malformations. Eur Radiol 2005;15:267–78
11. Basser PJ, Pierpaoli C. Microstructural and physiological features of tissues elucidated by quantitative-diffusion-tensor MRI. J Magn Reson B 1996;111:209–19
12. Basser PJ, Mattiello J, LeBihan D. Estimation of the effective self-diffusion tensor from the NMR spin echo. J Magn Reson B 1994;103:247–54
13. Rodesch G, Lasjaunias P. Spinal cord arteriovenous shunts: from imaging to management. Eur J Radiol 2003;46:221–32
14. Rodesch G, Hurth M, Alvarez H, et al. Angio-architecture of spinal cord arteriovenous shunts at presentation. Clinical correlations in adults and children. The Bicetre experience on 155 consecutive patients seen between 1981–1999. Acta Neurochir (Wien) 2004;146:217–26; discussion 226–27
15. Kringis T, Reinges MHT, Thiers R, et al. Functional and diffusion weighted magnetic resonance imaging of space-occupying lesions affecting the motor system: imaging the motor cortex and pyramidal tracts. J Neurol 2001;248:816–24
16. Reinges MHT, Schloth F, Coenen VA, et al. Imaging of postthalamalic visual fiber tracts by anisotropic diffusion weighted MRI and diffusion tensor imaging. Principles and applications. Eur J Radiol 2004;49:91–104
17. McCormick PC, Torres R, Post KD, et al. Intramedullary ependymoma of the spinal cord. J Neurol 1996;272:523–32
18. Basser PJ, Pierpaoli C. A simplified method to measure the diffusion tensor from seven MR images. Magn Reson Med 1998;39:928–34
19. Pierpaoli C, Basser PJ. Toward a quantitative assessment of diffusion anisotropy. Magn Reson Med 1996;36:903–906
20. Mori S, Crain B, Chacko V, et al. Three-dimensional tracking of axonal projections in the brain by magnetic resonance imaging. Ann Neurol 1999;45:265–69
21. Xu D, Mori S, Solaiyappan M, et al. A framework for callus fiber distribution analysis. NeuroImage 2002;17:1131–43
22. Westin CF, Maier SE, Mamata H, et al. Processing and visualization for diffusion tensor MRI. Med Image Anal 2002;6:93–108
23. Doppman JL, Di Ciero G, Dwyer AJ, et al. Magnetic resonance imaging of spinal arteriovenous malformations. J Neurol 1987;66:830–40
24. Dormont D, Assouline E, Gelbert F, et al. MRI study of spinal arteriovenous malformations. J Neuroradiol 1987;14:551–64
25. Rodesch G, Hurth M, Alvarez H, et al. Embolization of spinal cord arteriovenous shunts: morphological and clinical follow-up and results–review of 69 consecutive cases. Neuroradiology 2003;53:40–49; discussion 49–50
26. Maruyama K, Kamada K, Shin M, et al. Integration of three-dimensional corticospinal tractography into treatment planning for gamma knife surgery. J Neurosurg 2005;102:673–77
27. Beaulieu C, Allen PS. Determinants of anisotropic water diffusion in nerves. Magn Reson Med 1994;31:394–400
28. Elshafiey I, Bilgen M, He R, et al. In vivo diffusion tensor imaging of rat spinal cord at 7 T. Magn Reson Imaging 2002;20:233–47
29. Thonnard G, Glauche V, Weiller C, et al. Time course of wallerian degeneration after ischaemic stroke revealed by diffusion tensor imaging. J Neurol Neurosurg Psychiatry 2005;76:266–68
30. Werring DJ, Toosy AT, Clark CA, et al. Diffusion tensor imaging can detect and quantify corticospinal tract degeneration after stroke. J Neurol Neurosurg Psychiatry 2000;69:269–72
31. Voss HU, Watts R, Ulag AM, et al. Fiber tracking in the cervical spine and inferior brain regions with reversed gradient diffusion tensor imaging. Magn Reson Imaging 2006;24:231–39
32. Wang FN, Huang TY, Lin FH, et al. PROPELLER EPI: an MRI technique suitable for diffusion tensor imaging at high field strength with reduced geometric distortions. Magn Reson Med 2005;54:1232–40