Recent advances in magnetic resonance neuroimaging of lumbar nerve to clinical applications: A review of clinical studies utilizing Diffusion Tensor Imaging and Diffusion-weighted magnetic resonance neurography

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Abstract:

Much progress has been made in neuroimaging with Magnetic Resonance neurography and Diffusion Tensor Imaging (DTI) owing to higher magnetic fields and improvements in pulse sequence technology. Reports on lumbar nerve DTI have also increased considerably.

Many studies have shown that the use of DTI in lumbar nerve lesions, such as lumbar foraminal stenosis and lumbar disc herniation, makes it possible to capture images of interruptions of tractography at stenotic sites, enabling the diagnosis of stenosis. DTI can also reveal significant decreases in fractional anisotropy (FA) with significant increases in apparent diffusion coefficient (ADC) values in compression lesions.

FA values have higher accuracy than ADC values. Furthermore, strong correlations exist between FA values and indications of neurological severity, including the Japanese Orthopaedic Association (JOA) score, the Oswestry Disability Index (ODI), and the Roland-Morris Disability Questionnaire (RDQ) in patients with lumbar disc herniation-induced radiculopathy.

Most lumbar DTI has become 3T; 3T MRI has made it possible to take high-resolution DTI measurements in a short period of time. However, increased motion artifacts in the magnetic susceptibility effect lead to signal irregularities and image distortion. In the future, high-resolution DTI with reduced field-of-view may become useful in clinical applications, since visualization of nerve lesions and quantification of DTI parameters could allow more accurate diagnoses of lumbar nerve dysfunctions. Future translational studies will be necessary to successfully bring MR neuroimaging of lumbar nerve into clinical use.

Keywords:

magnetic resonance imaging, diffusion tensor imaging, diffusion-weighted MR neurography, lumbar nerve, lumbar foraminal stenosis, lumbar disc herniation

Introduction

With the rapid aging of our society, the number of patients with spinal disorders continues to rise. In the United States alone, 25 million patients complain of lumbar pain, leading to medical costs of over 100 billion dollars annually. Intervertebral disc lesions and lumbar nerve root disorders often cause lower back pain. Pain signals are transmitted from the local site to the peripheral nerves, via the spinal cord to the brain, where they are recognized as pain. In re-
cent years, functional MRI (fMRI) \cite{22} and MR spectroscopy \cite{23} have commonly been employed as neurofunctional imaging methods that involve the brain.

Lumbar nerve entrapment causes low back pain and radiculopathy, but discrepancies are often found between clinical symptoms and the degree of nerve root compression on conventional MR images. For instance, disc abnormalities are frequently observed on images from asymptomatic patients \cite{39}, so it has not been possible to understand physiologically the cause of the pain and to provide a quantitative evaluation of nerve damage.

Higher magnetic fields and improvements in pulse sequence technology in recent years have allowed for higher-resolution neuroimaging. Conventional 2-dimensional (2D) MR imaging cannot visualize the nerve roots in 1 image, because they cannot be depicted in any optional cross section.

3-dimensional MR neurography is useful for evaluating entire nerves, because it achieves better contrast between nerves and surrounding tissues. Recently, the use of MR neurography has increased for the evaluation of patients with suspected lumbosacral plexus involvement and for aiding in the confirmation of the diagnosis or providing anatomic information should surgical intervention be necessary \cite{65}.

However, this method could only demonstrate morphological abnormalities, such as nerve disruption and narrowing, but it could not quantitatively evaluate nerve damage.

Diffusion weighted imaging (DWI) and diffusion tensor imaging (DTI) are newer MR imaging techniques that measure the in vivo directional coherence of water diffusion. These quantitative techniques can identify pathology that cannot be detected by conventional MR imaging and thus, are important not only for research but also for clinical applications.

This article addresses developments in MR neuroimaging, such as diffusion-weighted MR neurography and DTI of the lumbar nerve in clinical applications.

### 1. Trends in clinical MR neuroimaging of lumbar nerves

We performed a structured search of the PubMed literature database to include relevant articles from 2007 to 2016. The following combinations of search terms were used: “lumbar nerve or lumbosacral plexus and diffusion-weighted MR neurography or DTI.” We excluded reviews, spinal cord studies and animal or in vivo studies. Our review identified a total of 35 articles: 24 DTI studies \cite{7-30} and 11 diffusion-weighted MR neurography studies \cite{14-15}. The number of lumbar nerve DTI studies has sharply increased in recent years (Fig. 1).

A systematic review of spinal cord MRI techniques, including DTI, magnetization transfer (MT), MRS, and fMRI, found that DTI is also being utilized more frequently and is the most mature among these tools \cite{22}.

### 2. Diffusion-weighted MR neurography

DWI based on MRI yields useful information regarding tissue microstructure because it monitors the random movement of water molecules, which is restricted by applying a motion probing gradient (MPG) in some directions \cite{44-46}. DWI data can be used to determine quantitative values, such as the apparent diffusion coefficient (ADC). DWI is frequently used to diagnose central nervous system diseases, such as acute stroke \cite{57}. It is also useful for the evaluation and diagnosis of lesions, such as those in multiple sclerosis \cite{48,49,52,53} and peripheral nerve compression disorders, such as carpal tunnel syndrome \cite{54}. Increased mean ADC values were observed in injured nerves with demyelination \cite{48,49,52,53}. Spinal cord imaging has technical limitations, such as the relatively small size of the cord, susceptibility artifacts from tissue-bone interfaces, and motion artifacts from respiratory activity \cite{56}. Yamashita and Takahara et al. \cite{57} (Fig. 2A) demonstrated the feasibility of whole-body MR neurography with DWI that can visualize tissues with impeded diffusion, such as tumors, the brain, the spinal cord, and the peripheral nerves \cite{56}. We found that diffusion-weighted MR neurography revealed that nerve swelling and transverse and mean ADC values were significantly greater in entrapped nerve roots in patients with lumbar disc herniation \cite{58} and lumbar foraminal stenosis (Fig. 2B, C) \cite{59}, suggesting demyelination and edema by slow compression caused by an increased degree of diffusion compared with that of normal tissue.

However, this method requires restructuring of 3D nerve images with maximum intensity projection (MIP) processing which is a volume rendering method for 3D imaging that projects the voxels with maximum intensity of the axial image, which leads to limitations in image resolution.
3. Diffusion Tensor Imaging (DTI)

DWI has a strong effect on not only the ease but also the directionality of water molecule dispersion. Axonal cell membranes and myelinated sheaths prevent dispersion in the direction directly aligned to the nerve fiber bundle, which leads to a loss of water molecule isotropy. This condition is referred to as anisotropy, and DTI and tractography are examples of selective recording of this data. FA is a parameter of anisotropic strength and is expressed in values from 0 to 1. The closer the value is to 1, the stronger the anisotropy, while a value of 0 is complete isotropy. DTI has recently been reported to be useful in demyelinating diseases, such as multiple sclerosis, or for chronic peripheral nerve compression lesions, such as carpal tunnel syndrome. Water molecule dispersion due to myelin sheaths in myelinated nerves in tissues occurs only in parallel to the nerve fibers, revealing high anisotropy, but when demyelination occurs as a result of nerve injuries such as spinal cord injury, the anisotropy decreases, leading to decreases in FA values.

3.1 Subjects of DTI lumbar nerve studies

Table 1 summarizes recent DTI lumbar nerve studies. A total of 683 subjects were studied across 24 DTI articles. 177 patients with lumbar spinal canal stenosis (LSCS) in 5 articles, 143 patients with lumbar disc herniation (LDH) in 7 articles, 112 patients with lumbar radiculopathy in 3 articles, 36 patients with foraminal stenosis (FS) in 6 articles, 2 patients with lower back pain in one article, 7 post mortem subjects were investigated in one article. 201 healthy volunteers were also investigated in 15 articles. 7 post mortem subjects were investigated in one article. 201 healthy volunteers were also investigated in 15 articles.

3.2 Measurement of DTI parameters

FA values were examined in all studies. All 17 articles demonstrated findings of low FA in the compressed nerves. ADC is the next most commonly reported parameter in 16 articles. Except for one

Figure 2. Whole-body MR neurography in coronal (left) and sagittal (right) views in a healthy volunteer (A). Taken from Yamashita et al. with permission from the Massachusetts Medical Society. Parasagittal T1-weighted image (B) and coronal neurography using b=1000 image (C) of a lumbar nerve root in a 66 year old man with L5-S1 foraminal stenosis. The arrowhead shows L5 foraminal stenosis with loss of the perineural fat signal (B). The arrow shows the entrapped nerve shifted upward and ran transversely in the foramen (C).
Table 1. Summary of Lumbar Nerve DTI Studies.

<table>
<thead>
<tr>
<th>Authors, Journal</th>
<th>Year</th>
<th>MRI B0; Vendor; Coil</th>
<th>DTI acquisition</th>
<th>Subjects</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Li et al. Biomed Rep.</td>
<td>2016</td>
<td>3.0T, GE, 8-channel spine coil</td>
<td>b=600 s/mm², 15 directions, ss EPI</td>
<td>LDH: 45</td>
<td>FA decreased, ADC decreased in the compressed nerves.</td>
</tr>
<tr>
<td>Haakma et al. Forensic Sci Int.</td>
<td>2016</td>
<td>3.0T, Philips, 16-channel surface coil</td>
<td>b=800 s/mm², 15 directions, ss EPI</td>
<td>Post-mortem (PM) subject: 7, Living subjects: 6</td>
<td>PM subjects showed lower diffusivity values compared to living subjects and fiber tractography results were comparable.</td>
</tr>
<tr>
<td>Eguchi et al. Asian Spine J.</td>
<td>2016</td>
<td>1.5T, Philips, SENSE-Spine-coil</td>
<td>b=800 s/mm², 15 directions, EPI</td>
<td>FS: one</td>
<td>FA decreased, ADC increased in compressed nerve of patient with FS.</td>
</tr>
<tr>
<td>Chhabra et al. World J Radiol.</td>
<td>2016</td>
<td>3.0T, Siemens</td>
<td>b=800,1,000 s/mm², 12 directions, EPI</td>
<td>Lumbar radiculopathy: 10</td>
<td>Individual differences were observed among neuropathic appearing nerve (low FA and increased ADC).</td>
</tr>
<tr>
<td>Eguchi et al. Asian Spine J.</td>
<td>2016</td>
<td>1.5T, Philips, SENSE-Spine-coil</td>
<td>b=800 s/mm², 15 directions, EPI</td>
<td>LSCS: 9, FS: 7, Volunteers: 5</td>
<td>Low FA and high ADC were marked in the extraforaminal zone for foraminal stenosis. FA showed higher accuracy than ADC.</td>
</tr>
<tr>
<td>Chen et al. J Orthop Surg Res.</td>
<td>2016</td>
<td>3.0T, Philips, 8-channel spine coil</td>
<td>b=600 s/mm², 15 directions, EPI</td>
<td>LSCS: 114</td>
<td>DTI showed clear benefits in determining decompression levels of LSCS than MRI. FA of positive levels decreased.</td>
</tr>
<tr>
<td>Eguchi et al. Bone Joint J.</td>
<td>2016</td>
<td>1.5T, Philips, SENSE-Spine-coil</td>
<td>b=800 s/mm², 15 directions, EPI</td>
<td>LDH: 13</td>
<td>There were strong correlations between FA and indications of neurological severity including JOA score and RDQ.</td>
</tr>
<tr>
<td>Manoliu et al. Invest Radiol.</td>
<td>2016</td>
<td>3.0T, Siemens</td>
<td>b=700 s/mm², readout-segmented, selective-excitation EPI</td>
<td>Volunteers: 12</td>
<td>DTI acquisitions in the coronal plane produced images of higher quality than the standard images in the axial orientation.</td>
</tr>
<tr>
<td>Wu et al. Spine</td>
<td>2016</td>
<td>1.5T, GE, 6 elements phased array spine coil</td>
<td>b=900 s/mm², 15 directions, EPI</td>
<td>LDH: 40</td>
<td>FA decreased in the compressed nerves. A significant negative association was observed between FA and ODI and symptom duration.</td>
</tr>
<tr>
<td>Kanamoto et al. Spine</td>
<td>2016</td>
<td>1.5T, Philips, SENSE-Spine-coil</td>
<td>b=800 s/mm², 15 directions, EPI</td>
<td>LSCS: 10, Double-crush lesion: 5, Volunteers: 5</td>
<td>Low FA and high ADC values indicating widespread nerve damage ranging from the medial intraspinal zone to the extraforaminal zone were noted in double-crush lesions.</td>
</tr>
<tr>
<td>Hou et al. Neural Regen Res.</td>
<td>2015</td>
<td>1.5T, GE, 8-channel cardiac coil</td>
<td>b=400 s/mm², 12 directions, EPI</td>
<td>LSCS: 31, Volunteers: 20</td>
<td>DTI showed abnormalities such as thinning and distortion in 49% and abruption in 23%. FA decreased in the lumbar sacral spinal nerve roots of patients with LSCS.</td>
</tr>
<tr>
<td>Miyagi et al. BMC Musculoskelet Disord.</td>
<td>2015</td>
<td>3.0T, Philips, 5-channel surface coil</td>
<td>b=800 s/mm², 33 directions, EPI</td>
<td>Volunteers: 6</td>
<td>FA increased and ADC decreased linearly up to 15 mm from the dura junction in the normal lumbar nerve roots.</td>
</tr>
<tr>
<td>Shi et al. Eur J Radiol.</td>
<td>2015</td>
<td>1.5T, GE, 8-channel cardiac coil</td>
<td>b=800 s/mm², 11 directions, EPI</td>
<td>Sciatica patients: 75, Volunteers: 36</td>
<td>FA decreased in compressed nerves. FA values were more sensitive and specific than MR imaging for differentiating compressed nerve roots, especially in the far lateral zone.</td>
</tr>
<tr>
<td>Oikawa et al. Magn Reson Imaging.</td>
<td>2015</td>
<td>3.0T, GE, sense XL Torso coil</td>
<td>b=800 s/mm², 11 directions, EPI</td>
<td>FS: 14, LSCS: 12, LDH: 5, low back pain: 2, spondylolisthesis: one</td>
<td>FA decreases in symptomatic roots. Abnormalities of tractography were found in patients with LSCS, and especially in patients with FS.</td>
</tr>
<tr>
<td>Sakai et al. J Med Invest.</td>
<td>2014</td>
<td>3.0T, Philips, 5-channel surface coil</td>
<td>b=800 s/mm², 33 directions, EPI</td>
<td>Asymptomatic LDH: one</td>
<td>In asymptomatic case, FA increased and ADC decreased following compression of the nerve root without injury.</td>
</tr>
<tr>
<td>Chuanting et al. Acta Radiol.</td>
<td>2014</td>
<td>3.0T, Philips, 15-channel spine coil</td>
<td>b=800 s/mm², 32 directions, EPI</td>
<td>LDH: 20, Volunteers: 20</td>
<td>FA decreased, MD increased in compressed nerve. Lumbar sacral root compression sites could be clearly identified on the tractography.</td>
</tr>
<tr>
<td>Karampinos et al. NMR Biomed.</td>
<td>2013</td>
<td>3.0T, GE, 6-cannel spine coil</td>
<td>b=500 s/mm², 24 directions, ss EPI</td>
<td>Volunteers: 11</td>
<td>rFOV method could minimize partial volume effect, breathing artifact and geometric distortion.</td>
</tr>
</tbody>
</table>
### Table 1

<table>
<thead>
<tr>
<th>Authors</th>
<th>Journal</th>
<th>Year</th>
<th>MRI B0; Vendor; Coil</th>
<th>DTI acquisition</th>
<th>Subjects</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Burdzik et al.</td>
<td>Eur Radiol.</td>
<td>2013</td>
<td>3.0T, Philips, cardiac coil</td>
<td>b=700 s/mm², 15 directions, s.s EPI</td>
<td>15 volunteers</td>
<td>FA decreased in compressed nerve of patient with FS.</td>
</tr>
<tr>
<td>van der Jagt et al.</td>
<td>Spine</td>
<td>2013</td>
<td>3.0T, Philips, cardiac coil</td>
<td>b=700 s/mm², 15 directions, s.s EPI</td>
<td>16 volunteers</td>
<td>FA decreased in compressed nerve of patient with FS.</td>
</tr>
<tr>
<td>Kimura et al.</td>
<td>Eur Radiol.</td>
<td>2011</td>
<td>3.0T, GE, 16-channel surface coil</td>
<td>b=700 s/mm², 11 directions, s.s EPI</td>
<td>8 volunteers</td>
<td>FA decreased in compressed nerve of patient with FS.</td>
</tr>
<tr>
<td>Eguchi et al.</td>
<td>AJNR</td>
<td>2011</td>
<td>3.0T, Philips, sense XL torso coil</td>
<td>b=800 s/mm², 11 directions, s.s EPI</td>
<td>19 volunteers</td>
<td>FA decreased, MD increased in compressed nerve.</td>
</tr>
<tr>
<td>Akindun et al.</td>
<td>Eur Radiol.</td>
<td>2010</td>
<td>1.5T, Philips, 8-channel spine coil</td>
<td>b=800 s/mm², 11 directions, s.s EPI</td>
<td>19 volunteers</td>
<td>FA decreased in the thoracic cords.</td>
</tr>
<tr>
<td>Filippi et al.</td>
<td>Eur Radiol.</td>
<td>2010</td>
<td>3.0T, GE, 15-channel spine coil</td>
<td>b=800 s/mm², 11 directions, s.s EPI</td>
<td>6 volunteers</td>
<td>FA decreased, MD increased in compressed nerve.</td>
</tr>
<tr>
<td>Chi et al.</td>
<td>Spine</td>
<td>2011</td>
<td>1.5T, GE, 15-channel spine coil</td>
<td>b=800 s/mm², 11 directions, s.s EPI</td>
<td>19 volunteers</td>
<td>FA decreased in compressed nerve of patient with FS.</td>
</tr>
<tr>
<td>Macnab et al.</td>
<td>Spine</td>
<td>2010</td>
<td>3.0T, GE, 16-channel surface coil</td>
<td>b=700 s/mm², 15 directions, s.s EPI</td>
<td>15 volunteers</td>
<td>FA decreased in compressed nerve of patient with FS.</td>
</tr>
<tr>
<td>Neugebauer et al.</td>
<td>Spine</td>
<td>2010</td>
<td>3.0T, Philips, 16-channel surface coil</td>
<td>b=800 s/mm², 15 directions, s.s EPI</td>
<td>15 volunteers</td>
<td>FA decreased in compressed nerve of patient with FS.</td>
</tr>
<tr>
<td>Macnab et al.</td>
<td>Spine</td>
<td>2009</td>
<td>3.0T, Philips, 15-channel spine coil</td>
<td>b=800 s/mm², 11 directions, s.s EPI</td>
<td>15 volunteers</td>
<td>FA decreased in compressed nerve of patient with FS.</td>
</tr>
</tbody>
</table>

### 3.3 DTI acquisition parameters

Parameters including magnetic field strength, motion-probing gradient (MPG), and b-values are important in the clinical use of DTI. High magnetic field strengths have the advantages of greater signal-to-noise ratio, improved spatial resolution, and faster imaging times, but also have the disadvantage of the magnetic susceptibility artifact. Magnetic susceptibility artifact indicated MRI artifacts that share distortions due to local inhomogeneous magnetic field. Fifteen reports used 3T, and 9 used 1.5T, so it appears that 3T research accounted for the majority.

The methods of measuring in vivo diffusion by MRI have been based mainly on the addition of several MPGs to the spin echo sequence to produce signal attenuation for the spins moving at random. Additional MPG directions can increase the resolution, but will prolong the scan time. MPG directions ranged from 11 to 33, where 15 directions was most common, used in 10 reports.

The b-value is a factor that defines gradient strength and duration and determines the degree of diffusion weighted images typically using values from 0 to 1000 s/mm². The higher the b-value, the stronger diffusion effects. Increasing the b-value increases the diffusion sensitivity, but decreases the signal-to-noise ratio. B-values ranged between 400 and 900, with b=800 the most common, used in 14 reports (Table 1).

### 3.4 Diagnosis of lumbar foraminal stenosis with DTI

In patients with lumbar foraminal stenosis, a nerve root or spinal nerve is entrapped in a narrowed lumbar foramen as the result of a degenerative lumbar spinal disorder. At this site, a dorsal root ganglion functions as a pain receptor; thus, the condition is refractory and may cause severe lower limb pain60. Macnab et al.60 appropriately named lumbar foraminal stenosis the “hidden zone,” as it is often overlooked, it accounts for approximately 60% of failed back surgery syndromes, and it decreases surgical success rates60.

Diagnostic imaging of lumbar spinal canal stenosis includes X-rays, CT, and MRI60-64, along with functional diagnosis via selective nerve root imaging and infiltration65. Conventional MRI has a false positive rate of 30% to 40%
in lumbar foraminal stenosis cases; thus, diagnosis is difficult. Recently 3D-CT, MR myelography, and 3D-MRI were reported to be useful for diagnosis.

We assessed the usefulness of DTI for the diagnosis of lumbar foraminal stenosis. Subjects in a supine position were scanned with a 3T MR imaging scanner (Discovery MR750; GE Healthcare, Milwaukee, WI) using a Sense XL Torso coil. We performed DTI with echo-planar imaging at a B value of 800 s/mm$^2$ and with 11 directional MPGs. Mean FA values and mean ADC values in the lumbar nerve roots were quantified on DTI images, and the lumbar nerve roots were visualized with tractography. In all subjects, the lumbar nerve roots were clearly visualized (Fig. 3A), and tractography also showed abnormalities, such as tract disruption, nerve narrowing, and indentation in their course through the foramen (Fig. 3B). Mean FA values were significantly lower, and mean ADC values were significantly higher, in entrapped versus intact roots.

We evaluated the accuracy of these parameters for the diagnosis of foraminal stenosis. Changes in DTI parameters (low FA and high ADC values) were marked in nerve roots and in the extraforaminal zone in foraminal stenosis. Additionally, the FA value was more accurate than the ADC value in the extraforaminal zone, and a low FA value suggested foraminal stenosis.

L5 radiculopathy may occur when the nerve is compressed at two levels (medial and lateral), for example, when the L4/S level is compressed by an intraspinal canal lesion and the L5/S1 level is compressed by a lateral lesion. This is called a double-crush lesion. Traditional imaging studies do not allow a determination of whether the compressing lesion is inside or outside of the spinal canal, or if a double-crush lesion is responsible, which leads to poor surgical success rates. The distal latency of L5 has been measured electrophysiologically; however, this is an invasive approach, and noninvasive diagnostic methods are virtually nonexistent. Diagnosis of L5 nerve damage double-crush lesions by DTI was studied. Low FA values and high ADC values indicative of widespread nerve damage from the medial intraspinal zone to the extraforaminal zone were found for double-crush lesions. If double-crush injury is suspected prior to surgery, DTI assessment may help prevent failed back surgery syndrome.

3.5 DTI evaluation of radiculopathy in patients with lumbar disc herniation

Mixter and Barr first described radicular pain of the sciatica as spinal root compression by a herniated intervertebral disc; however, the underlying pathophysiology is not well understood. In clinical practice, asymptomatic intervertebral disc degeneration and herniation are often found, and these discrepancies can confuse spine surgeons. We previously reported a correlation between neurological severity and DTI parameters, such as the FA and ADC values, in patients with radiculopathy caused by lumbar disc herniation. Strong correlations were found between the FA value and indications of neurological severity, including the Japanese Orthopedic Association (JOA) score, the Oswestry Disability In-
Figure 4. Presurgical lumbar MRI images and tractography of an 87 year old woman with double-crush lesions. She complained of persistent symptoms due to L5 foraminal stenosis despite L4/5 decompression surgery. A) Sagittal T2-weighted image B) Right parasagittal T1-weighted image. Right L5 foraminal stenosis (arrow) can be noted. C) L3/4 level axial image D) Coronal 3D-MR image. Right L5 nerve swelling and running transversely in the course through the foramen (arrowhead). E) Fusion image of lumbar DTI images and 3D-T2 weighted images. The right L5 nerve is disrupted at the foraminal area (arrowhead). F) FA values (intraspinal zone, nerve root zone, and extraforaminal zone) were 0.301, 0.375, and 0.361, respectively, on the affected side and 0.392, 0.416, and 0.434, respectively, on the unaffected side. FA values were decreased over a widespread area from the intraspinal to the extraforaminal zone.

Figure 5. Tractography of lumbar nerves in a patient with lumbar disc herniation between the L5 and S1 discs before and after microendoscopic discectomy (MED). Tractography of the S1 nerve on the right side was disrupted by disc herniation (arrow) before MED. At six months after MED, leg numbness was decreased from 60 to 10, the FA value was increased from 0.299 to 0.327, and the ADC value was decreased from 1.173 mm$^2$/s to 1.096 mm$^2$/s. Tractography of the S1 nerve was elongated to the proximal side (arrowheads).
Figure 6. Tractography of the L5 and S1 nerves in a healthy volunteer with reduced FOV (rFOV) (A) and conventional FOV (cFOV) (B). Axial FA map at the L4/5 level in a healthy volunteer with rFOV (C) and cFOV (D). Fiber counts are higher with rFOV versus cFOV, allowing for clearer imaging of the lumbar nerve. FA map resolution is higher with rFOV allowing for clearer imaging of the nerve root (arrow head) and the spinal canal (arrow).

dex (ODI)\textsuperscript{15}, and the Roland-Morris Disability Questionnaire (RDQ)\textsuperscript{13}. DTI parameters are useful for diagnosis, quantitative assessment, and follow-up of lumbar nerve entrapment (Fig. 5).

Advantages and limitations of DTI

It has not been possible to understand the physiological cause of pain by conventional MR images because of the discrepancy between clinical symptoms and MRI findings, which may be an issue for diagnostic and therapeutic management. Conventional MR imaging also cannot selectively visualize peripheral nerves and cannot quantitatively assess the severity of the nerve lesion. DTI is a non-invasive way to effectively trace the nerve fiber bundle, and quantitatively evaluate the nerve injury.

DTI has several limitations. Firstly, tractography is mathematical modeling of the diffusion tensor data using probability theories to model the most likely course of diffusion, and the number of tracts visualized by DTI did not present the actual volume of nerve fiber trajectories. Second, 3T MRI has made it possible to take high-resolution DTI measurements in a short period of time. However, since it is a high magnetic field, the magnetic susceptibility effect and increases in motion artifacts lead to signal irregularities and image distortion so that nerve fiber follow-up is limited in areas with artifacts. Third, the evidence is insufficient to support DTI as a diagnostic tool or predictor of clinical outcomes. Fourth, automatic analysis methods such as tract-specific automatic ROI placement are needed.

Further translational research is required to link pathophysiologic mechanisms in animal models to clinical outcomes in patients.

3.6 Future directions; High-resolution DTI

In the future, further improvements in image resolution are essential for clinical applications. A new reduced field-of-view FOV (rFOV) single-shot diffusion-weighted echo-planar imaging method that uses a 2D spatially selective echo-planar RF excitation pulse and a 180° refocusing pulse to reduce the FOV in the phase-encode (PE) direction, and simultaneously suppresses the signal from fat has been proposed. The rFOV method decreases the readout duration and allows acquisition of high-resolution diffusion-weighted images for practical application to spinal imaging\textsuperscript{71}.

rFOV DTI of the spinal cord allows for acquisition of high-resolution images for assessment of specific spinal cord tracts and discrimination of white matter from gray matter. Maki et al.\textsuperscript{72} showed correlations between the FA of specific white matter in spinal cord tracts and myelopathy severity in
patients with cervical compression myelopathy with rFOV DTI. Several authors\textsuperscript{4,25} reported that rFOV imaging was necessary for lumbar nerve root DTI imaging and tractography in 3T, as it drastically reduced the susceptibility and chemical shift artifacts that hindered accurate tractography on conventional FOV imaging.

We attempted high-resolution imaging of the lumbar nerves with rFOV 3T MRI. Compared to traditional methods, rFOV allows for clear imaging of the lumbar nerves and enables accurate measurements of the FA and ADC values (Fig. 6A-D). By expressing consecutive FA changes in color, we found that it was possible to visualize the degree of nerve damage (Fig. 7). This method allows for direct visualization of consecutive changes in FA values, and it may break through the limitations of ROI establishment such as the partial volume effect.

In the future, high-resolution DTI with rFOV may be used to visualize nerve lesions and allow for more accurate diagnosis of DTI parameter quantification with opportunities for clinical applications.

**Conclusions**

Recent advances have allowed the use of higher magnetic fields and pulse sequencing improvements in MRI devices that provide better neuroimaging, such as DTI in the lumbar nerves. DTI of spinal nerves is more likely to be affected by the magnetic rate than the brain; therefore, from a technical viewpoint, it can hardly be considered a widely employed test in the field, but reports continue to suggest that it may reveal information that is not available on a conventional MRI.

In the future, introduction of high-resolution DTI will allow more detailed visualization of lumbar nerve lesions that may be quantified, and we look forward to further expansion in the functional diagnosis of lumbar nerve disorders. Future translational studies will be essential to successfully bring these quantitative techniques into clinical use.

Conflicts of Interest: The authors declare that there are no conflicts of interest.

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