Reduced Field-of-View Diffusion Imaging of the Human Spinal Cord: Comparison with Conventional Single-Shot Echo-Planar Imaging

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BACKGROUND AND PURPOSE: DWI of the spinal cord is challenging because of its small size and artifacts associated with the most commonly used clinical imaging method, SS-EPI. We evaluated the performance of rFOV spinal cord DWI and compared it with the routine fFOV SS-EPI in a clinical population.

MATERIALS AND METHODS: Thirty-six clinical patients underwent 1.5T MR imaging examination that included rFOV SS-EPI DWI of the cervical spinal cord as well as 2 comparison diffusion sequences: fFOV SS-EPI DWI normalized for either image readout time (low-resolution fFOV) or spatial resolution (high-resolution fFOV). ADC maps were created and compared between the methods by using single-factor analysis of variance. Two neuroradiologists blinded to sequence type rated the 3 DWI methods, based on susceptibility artifacts, perceived spatial resolution, signal intensity–to-noise ratio, anatomic detail, and clinical utility.

RESULTS: ADC values for the rFOV and both fFOV sequences were not statistically different (rFOV: 1.01 ± 0.18 × 10⁻³ mm²/s; low-resolution fFOV: 1.12 ± 0.22 × 10⁻³ mm²/s; high-resolution fFOV: 1.10 ± 0.21 × 10⁻³ mm²/s; F = 2.747, P > .05). The neuroradiologist reviewers rated the rFOV diffusion images superior in terms of all assessed measures (P < 0.0001). Particular improvements were noted in patients with metal hardware, degenerative disease, or both.

CONCLUSIONS: rFOV DWI of the spinal cord overcomes many of the problems associated with conventional fFOV SS-EPI and is feasible in a clinical population. From a clinical standpoint, images were deemed superior to those created by using standard fFOV methods.

ABBREVIATIONS: ADC = apparent diffusion coefficient; AP = anteroposterior; DTI = diffusion tensor imaging; DWI = diffusion-weighted MR imaging; EPI = echo-planar imaging; fFOV = full FOV; HiRes, high resolution; isoDWI = isotropic DWI; LoRes, low resolution; RF = radio frequency; rFOV = reduced FOV; ROI = region of interest; SNR = signal intensity to noise ratio; SS-EPI = single-shot EPI; STIR = short tau inversion recovery; ZOOM-EPI = zonal oblique multisection EPI

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DWI and compare it with the routine fFOV SS-EPI method in a clinical population.

Materials and Methods

Industrial Support

The pulse sequence used is a collaborative “work-in-progress” between Stanford University and General Electric Healthcare.

Patient Population

MR imaging studies were acquired as part of routine clinical care. All patients provided written informed consent approved by our institutional review board for a prospective study of the safety and utility of advanced imaging sequences between March and July 2009. Patients with cervical spine examinations performed at 1.5T for clinical purposes were enrolled in the study. Three different magnets (Signa, GE Healthcare, Waukesha, Wisconsin; 40 mT/m maximum gradient strength and 150 mT/m/ms maximum slew rate) were used. Thirty-six patients (18 males, 18 females; mean age, 52 ± 19 years; range, 17–85 years) met the entry criteria. Imaging indications were similar to those that are routine at our institution (posttrauma, n = 11; neck pain, n = 7; known or suspected neoplasm, n = 4; weakness, n = 3; sensory abnormalities, n = 3; combined weakness and sensory abnormalities, n = 3; prior examination with incidental finding, n = 2; neurodegenerative disease, n = 1; known or suspected infection, n = 1; postsurgical follow-up, n = 1).

Imaging Methods

For all studies, the built-in body volume coil was used for signal intensity transmission and an 8-channel cervical-thoracic-lumbar coil for signal intensity reception. All patients received standard anatomic imaging of the cervical spine, which at our institution includes sagittal T1-weighted (TR/TE, 2400/11 ms), sagittal T2-weighted fast spin-echo (TR/TE, 500/22 ms), sagittal T2-weighted fast spin-echo (TR/TE, 4615/50/140 ms), gradient-echo (TR/TE, 475/11 ms) images.

The diffusion-weighted spin-echo rFOV SS-EPI images were acquired in the sagittal plane, by using six 4-mm sections with no intersection gap. The rFOV was achieved by a 90° 2D echo-planar RF excitation followed by a refocusing 180° pulse. A full description of the method can be found in Saritas et al., and a pulse sequence diagram is included in On-line Fig 1. TR/TE/EPI readout time was 3600/64/64 ms; FOV, 16 × 4 cm²; TR, 3602 ms; and matrix, 96 × 96. In the first patient, an additional fFOV scan was performed by using the following parameters: TR/TE/EPI, 4800/128/128 ms; FOV, 128 × 128 cm²; and matrix, 128 × 128.

Table 1: Imaging parameters for the DWI sequences

|              | rFOV | LoRes fFOV | HiRes fFOV |
|--------------|------|------------|------------|
| TR (ms)      | 3600 | 3600       | 3600       |
| TE (ms)      | 98   | 98         | 98         |
| FOV (cm²)    | 4.0  | 4.0        | 4.0        |
| Matrix size  | 192  | 192        | 192        |
| Bandwidth    | 62.5 | 62.5       | 62.5       |
| Readout (ms) | 54   | 59         | 216        |
| In-plane res. | 0.9  | 0.9        | 0.9        |
| Section thk. | 4    | 4          | 4          |
| Acquisition time (min:s) | 2:30 | 2:30       | 2:30       |

Radiologic Assessment

IsoDWI and ADC images from the 3 different DWI sequences were blinded and placed in random order by using a Fisher-Yates shuffle. Two neuroradiologists, each with >15 years of academic experience (C.T.C. and N.J.F.), evaluated each of the sequences (rFOV, low-resolution fFOV, and high-resolution fFOV) on the following imaging characteristics: anatomic detail, susceptibility and/or susceptibility-induced artifacts; perceived spatial resolution; perceived SNR; and perceived clinical utility. Scoring was performed based a 5-point scale as follows: 1 = nondiagnostic, 2 = poor, 3 = satisfactory, 4 = good, and 5 = excellent.

Another neuroradiologist (J.B.A.) manually outlined circular ROIs on the midline sagittal image with an area and volume of 0.19 cm² and 0.07 cm³, respectively, to compare ADC values among the different sequences. This size was chosen to be small enough to be placed within the cervical cord on all images regardless of field inhomogeneity-induced distortion, which often narrowed the apparent cord diameter on fFOV images, but to be large enough to obtain a reasonable number of voxels with which to calculate ADC (22 voxels for the rFOV and high-resolution fFOV and 5 voxels for the low-resolution fFOV). ROIs were placed immediately posterior to the inferior aspect of the C2 vertebral body such that the ROI was tangent and superior to a line drawn parallel to and passing through the C2–3

Table 1: Imaging parameters for the DWI sequences

|              | rFOV | LoRes fFOV | HiRes fFOV |
|--------------|------|------------|------------|
| TR/TE (ms)   | 3600 | 3600       | 3600       |
| FOV (cm²)    | 4.0  | 4.0        | 4.0        |
| Matrix size  | 192  | 192        | 192        |
| Bandwidth    | 62.5 | 62.5       | 62.5       |
| Readout (ms) | 54   | 59         | 216        |
| In-plane res. | 0.9  | 0.9        | 0.9        |
| Section thk. | 4    | 4          | 4          |
| Acquisition time (min:s) | 2:30 | 2:30       | 2:30       |

Although not part of the comparison study, 3 additional patients had rFOV imaging performed and findings are presented in the online material as examples of axial DWI of the cervical cord and sagittal thoracic spinal cord diffusion in patients with and without pathology. Axial imaging of the cervical spine was performed in a 53-year-old man with a history of trauma (On-line Fig 2). Parameters for this scan were as follows: TR/TE, 3600/64; FOV, 16 × 4 cm²; matrix, 192 × 48; voxel size, 0.83 cm²; and 6-mm section thickness with 1.5-mm skip. Thoracic spinal cord imaging is presented in 2 patients: a 49-year-old man scanned to evaluate known nerve sheath tumors (On-line Fig 3); and an 57-year-old man with known type B aortic dissection, and more recent history of acute onset paraparesis and chest pain radiating to back (On-line Fig 4). Sagittal rFOV imaging with the following parameters was performed in both of these patients: TR/TE; FOV, 30 × 6 cm; matrix, 320 × 64; and voxel size, 0.94 cm². In the first patient, an additional fFOV scan was performed by using the following parameters: TR/TE, 3000/69; FOV, 30 cm²; matrix, 128 × 128; and voxel size, 2.3 cm².
disk space. This location was chosen because it was relatively free of distortion and artifacts regardless of the acquisition technique, sequence, patient body habitus, or position within the scanner.

**Statistical Analysis**

All statistical analyses were performed by a biostatistician (J.R.) using Stata release 9.2 (StataCorp, College Station, Texas). ADC was compared among the 3 diffusion scans by using single-factor analysis of variance, and \( P < .05 \) was considered significant. Agreement between readers was assessed with weighted \( \kappa \) and exact Bowker test of symmetry. Pair-wise comparisons of neuroradiologist reviewers’ evaluations of the different DWI sequences were done with paired Wilcoxon tests.

**Results**

No abnormal findings, including reduced or increased diffusion, were noted in the patients included in this study, with the exception of varying degrees of degenerative disk disease. T2-weighted anatomic images, rFOV isoDWI, and ADC maps are shown for a representative case in Fig 1. We observed improvements in imaging artifacts by using rFOV isoDWI compared with our standard low-resolution fFOV isoDWI, particularly distortions in the phase-encode direction (AP on our images). An example of this is shown in Fig 2, in which the rFOV method has improved anatomic appearance compared with either of the fFOV methods. The fFOV images demonstrate pixel misregistration in the AP direction due to susceptibility. Also, artifacts seen at the edges of the disks and the CSF space of the spinal canal are significantly reduced on the rFOV images. Due to the prolonged readout with the high-resolution fFOV SS-EPI, T2*-induced blurring is also much stronger than on the rFOV scans with identical spatial resolution.

An example of axial imaging of the cervical spinal cord is shown as On-line Fig 2. Although fFOV diffusion images were not acquired in this patient, the general quality of the images are comparable or better than the typical scans acquired at our institution by using fFOV. For the purposes of example, a single comparison of rFOV and fFOV diffusion imaging of the thoracic cord is shown in On-line Fig 3. The higher resolution and reduced distortion has enabled the visualization of a small central CSF cavity in the distal thoracic cord, the terminal ventricle, which is not evident on the fFOV images. A further example of thoracic spinal cord diffusion imaging is shown in On-line Fig 4 that also demonstrates high signal intensity on DWI with associated low ADC in the central cord in the lower thoracic spine consistent with acute spinal cord infarct.

Figure 3 shows the location of the ROI in which the ADC measurements were made. The ADC values obtained with the rFOV approach were not statistically different from either of the fFOV diffusion techniques (rFOV, \( 1.01 \pm 0.18 \times 10^{-3}\) mm\(^2\)/s; low-resolution fFOV, \( 1.12 \pm 0.22 \times 10^{-3}\) mm\(^2\)/s; high-resolution fFOV, \( 1.10 \pm 0.21 \times 10^{-3}\) mm\(^2\)/s; \( F = 2.747, P > .05 \)). There is reduced interpatient variance for the rFOV method that may be related to the improved anatomic fidelity of the images.

Agreement between the 2 neuroradiologist reviewers was moderate to good (Table 2). The readers did not show a tendency to over- or underscore compared with each other, except for the category of anatomic detail, in where reviewer 2 gave higher ratings overall compared with reviewer 1 \( (P = .011) \). The reviewers preferred the rFOV images by a strong margin in all metrics (Fig 4). All 3 sequences were significantly different from one another on all ratings, with rFOV having consistently higher ratings than the low-resolution fFOV sequence, which in turn was preferred compared with the high-resolution fFOV sequence (all calculated \( P \) values < .0001) (Fig 4). This was most pronounced for the preference of the rFOV method on the basis of reduction of susceptibility artifacts, in which the rFOV images were scored good or excellent 69% of the time, compared with 7% for either of the fFOV images. Likewise, clinical utility was deemed excellent 51% of the time for the rFOV diffusion images versus 1% for either of the fFOV methods. Improvements in anatomic fidelity are illustrated by the case of a patient with left-sided arm weakness and multiple posterior disk-osteophyte complexes, the largest of which is at C4–5; the fFOV images demonstrate significant misregistration and image degradation in the regions of the osteophytes, which is largely alleviated by the rFOV approach (Fig 5). Another case of a patient with a C6–7 anterior cervical...
diskectomy and fusion demonstrates that the susceptibility artifact from the metallic plate is mitigated though not entirely eliminated by the rFOV method (Fig 6).

**Discussion**

This study demonstrates that a rFOV approach using a 2D RF pulse enables acquisition of high-resolution spinal cord diffusion images with reduced artifacts compared with conventional SS-EPI. By reducing the FOV, it is possible to obtain images at a given spatial resolution with a reduced number of $k$-space lines. For single-shot imaging, this directly translates to a shorter readout time and higher (pseudo) bandwidth along the phase-encode dimension; consequently, imaging artifacts such as blurring and pixel misregistration are reduced. The ability to acquire an asymmetric FOV is particularly well suited for sagittal imaging of the spinal cord, given that the relevant anatomic structures are elongated in the superior-inferior direction relative to the AP direction. The current study demonstrates that the combination of higher spatial resolution, decreased imaging artifacts, and good coverage of the relevant anatomy improves spinal cord DWI in clinical patients, as evaluated by 2 experienced neuroradiologists.

Improved anatomic depiction of the spinal cord by using rFOV was most notable in regions immediately ventral to the posterior spinous processes, which were often marred by a significant alteration of perceived AP cord diameter, cord signal intensity, and cord ADC values on the fFOV diffusion images (eg, Fig 5). Also, there were improvements in anatomic fidelity around metal (Fig 6) on the rFOV images. Both readers preferred the rFOV diffusion images to either of the fFOV images in all metrics by using a subjective 5-point scale. This preference was strongest for clinical utility, possibly because it represents the combined effect of the other assessed attributes.

![Fig 2](image1)

**Fig 2.** A 71-year-old woman with right-sided weakness and hemifacial spasm. Comparison of DWI by using either rFOV (A); fFOV matched to have the same readout time (with subsequently lower spatial resolution) (B); or fFOV matched to have the same spatial resolution, with increased readout time (and associated susceptibility artifacts) (C). All images were matched for acquisition time (2.5 minutes). Only the 4 central sections are shown. rFOV DWI demonstrates high spatial resolution with minimal artifacts and acceptable SNR.

![Fig 3](image2)

**Fig 3.** rFOV (A), low-resolution fFOV (B), and high-resolution fFOV ADC maps (C) demonstrating the location of the ROI used for the ADC measurements, placed on the centermost section of the sagittal images at the level of C2–3.

| Measure      | Weighted $\kappa$ | 95% CI       | Symmetry Test, $P$ Value |
|--------------|-------------------|--------------|--------------------------|
| Susceptibility | 0.58              | 0.48–0.66    | .968                     |
| Resolution   | 0.53              | 0.40–0.63    | .767                     |
| SNR          | 0.45              | 0.35–0.56    | .405                     |
| Anatomy      | 0.53              | 0.42–0.62    | .011                     |
| Clinical utility | 0.59            | 0.48–0.68    | .468                     |
such as improved susceptibility effects, SNR, and spatial resolution. Although still significant, the preference based on SNR of the resultant images was not as strong, with a significant number of the low-resolution fFOV images also deemed good or excellent. One must keep in mind, however, that the voxel size of the low-resolution fFOV images was 4 times larger than the rFOV images. High resolution is critical for imaging of the spinal cord given its small dimensions; when one compares the preference between the rFOV and fFOV images at fixed high spatial resolution, there is strong preference for the rFOV images (Fig 4).

ADC values were measured using the rFOV method in the upper cervical cord and were not statistically different from measurements made with conventional fFOV methods. Mean ADC values were within the range of measurements in previous reports.1,3,5,24,49,50

Several factors suggest that rFOV DWI may be a clinically feasible approach to image the spinal cord. The rFOV approach is time-efficient, as evidenced by the short duration (2.5 minutes) of the scans obtained in the current study. Although multishot interleaved methods would be expected to have similar SNR, the combination of data from multiple ac-

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**Fig 4.** Pooled reviewer ratings of anatomy (A), perceived resolution (B), perceived SNR (C), susceptibility artifact (D), and clinical utility (E) for each of the 3 assessed sequences: High-resolution fFOV (HiRes fFOV), low-resolution fFOV (LoRes fFOV), and rFOV. All 3 sequences were significantly different from each other on all ratings, with rFOV having consistently higher ratings than the LoRes fFOV sequence, which in turn was preferred over the HiRes fFOV sequence (all calculated P values ≤ .0001). All values are expressed as a percentage of total responses by both reviewers.
quisitions is less robust and overall imaging time is increased. Furthermore, because the modification to SS-EPI is only in the excitation pulse, if further improvements in susceptibility artifacts or higher spatial resolution are required, then it can be combined in a straightforward manner with other approaches, including parallel and multishot/interleaved EPI, though at a cost of longer scan time. Another benefit of single-shot acquisition is that by acquiring an entire section in a single excitation, and an entire contiguous volume within a TR interval, rFOV DWI is amenable to both 2D and 3D motion correction methods. Eddy current correction was not applied. This was because a lower $b$-value was used than in the brain. The higher effective bandwidth of rFOV also diminished geometric distortion effects emanating from eddy currents induced from switching diffusion-encoding gradients.

Another approach to acquire an rFOV is to suppress or dephase the signal intensity from tissue outside the desired FOV by using additional RF pulses. These techniques may be contrasted with the rFOV approach used in this study, which, rather than exciting and then suppressing/dephasing such regions, does not excite these regions in the first place. One such method, ZOOM-EPI, relies on a 180° refocusing pulse that is oblique to the initial section, such that only the parallelogram-shaped volume that experiences both the initial 90° and 180°
pulses contributes to the signal intensity during readout; this method was applied to image the spinal cord and the optic nerve, but it cannot be used to acquire contiguous sections without an SNR penalty, because the regions excited by the 2 pulses overlap with the adjacent sections. This is particularly problematic for imaging in the sagittal plane, because the entire spinal cord is only approximately 1 cm wide. Two other methods use refocusing pulses that are selective in the phase-encode direction to readout a rectangular FOV.44,46 The first method, known as contiguous ZOOM-EPI,46 uses 2 refocusing pulses such that spins outside the fFOV within contiguous sections are minimally affected; this also has the advantage of limiting eddy current distortions.51 Another method places section pulses to saturate the signal intensity outside the desired region,47 uses successive outer volume suppression pulses. This method has been applied to study diffusion in an SNR decrease due to T1 recovery between the 2 refocusing pulses overlap with the adjacent sections. However, the neighboring sections can still experience variations in the transmit B1 field. In summary, all of these considerations, because we wanted to acquire multiple diffusion sections would be possible with 2 acquisitions, requiring 5 minutes for the same SNR as the sagittal images. Also, we did not perform a comparison study of the different methods in the thoracic spine, where increased motion, including cardiac, respiratory, and CSF flow, is greater than in the cervical spine. However, our experience in a limited number of patients (eg, On-line Figs 3 and 4) suggests that rFOV performs well in the spinal cord. This may be due to the greater reductions in the phase-encode FOV due to more elongated contour of the thoracic spine in most individuals.

We chose to examine fFOV diffusion imaging with equivalent section thickness and imaging time. We did not compare with other competing approaches, including multishot interleaved EPI, parallel EPI, or outer volume suppression methods. Also, the fFOV images were acquired only by using a standard square FOV SS-EPI sequence. Even with a nonselective excitation pulse, it may be possible to read out a slightly rectangular FOV with minimal aliasing artifacts due to the geometry of the cervical spine. Placement of anterior saturation bands would also reduce aliasing and breathing artifacts with this approach. We did not choose to optimize the fFOV DWI scans in this manner, because this has not been part of our traditional clinical routine. Thus, the advantages of the rFOV approach that we describe may be less apparent if somewhat more optimized diffusion imaging methods are considered.

We did not perform DTI as part of this study; thus, we cannot report on longitudinal and parallel ADC values or fractional anisotropy. However, there is no inherent limitation in the algorithm for DTI,54 and it was omitted in the current study only because our goal was to assess overall image quality rather than these other metrics. Improved anatomic fidelity and high spatial resolution in the underlying diffusion images is critical for DTI studies in the spinal cord, and this topic will be a subject of future studies.

Finally, only relatively mild abnormalities were noted on these clinical studies, limited to degenerative disk disease; specifically, no diffusion abnormalities were observed on any of the studies. This is in keeping with the relatively uncommon nature of spinal cord infarction and other pathologies in the typical patient population referred for cervical spine examinations. However, our clinical experience suggests that the method aids in the detection of pathology, as demonstrated in the case of thoracic spinal cord infarction shown in On-line Fig 4. In future reports, we plan to describe in a more comprehensive manner the findings of rFOV DWI in a larger group of patients that includes such pathologies, but does not receive comparison fFOV SS-EPI, to allow us to comment on important issues such as lesion conspicuity.

Conclusions
Reduced FOV spinal cord DWI is feasible in a clinical population, improves image quality, and is preferred over conventional fFOV SS-EPI in a clinical population. Further studies to assess conspicuity and clinical utility for visualizing commonly encountered spinal cord pathologies are warranted.

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