Comparison of Sagittal FSE T2, STIR, and T1-Weighted Phase-Sensitive Inversion Recovery in the Detection of Spinal Cord Lesions in MS at 3T

P. Alcaide-Leon, A. Pauranik, L. Alshafai, S. Rawal, J. Oh, W. Montanera, G. Leung, and A. Bharatha

ABSTRACT

BACKGROUND AND PURPOSE: Determining the diagnostic accuracy of different MR sequences is essential to design MR imaging protocols. The purpose of the study was to compare 3T sagittal FSE T2, STIR, and T1-weighted phase-sensitive inversion recovery in the detection of spinal cord lesions in patients with suspected or definite MS.

MATERIALS AND METHODS: We performed a retrospective analysis of 38 patients with suspected or definite MS. Involvement of the cervical and thoracic cord segments was recorded on sagittal FSE T2, STIR, and T1-weighted phase-sensitive inversion recovery sequences independently by 2 readers. A consensus criterion standard read was performed with all sequences available. Sensitivity, specificity, and interobserver agreement were calculated for each sequence.

RESULTS: In the cervical cord, the sensitivity of T1-weighted phase-sensitive inversion recovery (96.2%) and STIR (89.6%) was significantly higher \( (P < .05) \) than that of FSE T2 (50.9%), but no significant difference was found between T1-weighted phase-sensitive inversion recovery and STIR. In the thoracic cord, sensitivity values were 93.8% for STIR, 71.9% for FSE T2, and 50.8% for T1-weighted phase-sensitive inversion recovery. Significant differences were found for all comparisons \( (P < .05) \). No differences were detected in specificity. Poor image quality and lower sensitivity of thoracic T1-weighted phase-sensitive inversion recovery compared with the other 2 sequences were associated with a thicker back fat pad.

CONCLUSIONS: The use of an additional sagittal sequence other than FSE T2 significantly increases the detection of cervical and thoracic spinal cord lesions in patients with MS at 3T. In the cervical segment, both STIR and T1-weighted phase-sensitive inversion recovery offer high sensitivity and specificity, whereas in the thoracic spine, STIR performs better than T1-weighted phase-sensitive inversion recovery, particularly in patients with a thick dorsal fat pad.

ABBREVIATION: PSIR = T1-weighted phase-sensitive inversion recovery

MR imaging of the spinal cord is an important diagnostic technique in MS because the prevalence of spinal cord abnormalities in patients with clinically isolated syndrome is as high as 42%.\(^1\) In clinically diagnosed MS, spinal cord involvement reaches 75%–92%, depending on the series.\(^2\)-\(^4\) The presence of asymptomatic cord lesions contributes to the demonstration of dissemination in space in the McDonald 2010 criteria for MS, and imaging of the spinal cord allows an increase of 18.3% in the number of patients meeting the diagnostic criteria.\(^5\) The presence of spinal cord lesions not only facilitates diagnosing MS but is also predictive of conversion to clinically definite MS, especially in patients with nonspinal clinically isolated syndrome who do not fulfill brain MR imaging criteria.\(^6\) Moreover, spinal cord lesions in MS can occur in isolation in 5% of patients, particularly in primary-progressive MS.\(^7\)

Spinal cord imaging is challenging because the spinal cord is a small and mobile structure.\(^8\) In addition, its anatomic location...
makes it prone to ghosting artifacts caused by the heart and great vessels as well as truncation artifacts. 3T MR imaging compared with 1.5T is more prone to artifacts caused by $B_1$ field inhomogeneity, susceptibility, vascular pulsation, and chemical shift.\textsuperscript{10,11} In addition, 3T MR imaging has a higher energy deposit within the tissue, resulting in a higher specific absorption rate than lower field scanners. These problems can be partially solved with various technical adjustments and fast (parallel) imaging.\textsuperscript{12}

Traditionally, the spinal cord in patients with MS has been imaged by using sagittal and axial FSE T2/proton density sequences. Additional sequences, including STIR\textsuperscript{13,14} and T1 inversion recovery,\textsuperscript{15} have shown promise by increasing lesion visibility, particularly at 3T, in which conventional FSE T2 and proton density images are frequently unsatisfactory.\textsuperscript{16} STIR has proved very useful as a complementary sequence in the detection of MS lesions but cannot be used in isolation due to its lower specificity.\textsuperscript{13,14} Numerous studies have demonstrated the superiority of STIR over T2 at 1.5T,\textsuperscript{13,14,17-19} and 1 study\textsuperscript{15} also showed the advantages of STIR at 3T in the cervical cord. To our knowledge, no studies have been performed in the thoracic cord comparing sagittal FSE T2 and STIR. A recent publication showed the advantages of T1-weighted phase-sensitive inversion recovery (PSIR) for the detection of cervical spinal cord lesions in MS at 3T.\textsuperscript{13} PSIR has been shown to improve lesion localization and boundary definition over STIR in the cervical spinal cord, but it has not been tested in the thoracic cord.\textsuperscript{16}

The aim of our study was to compare the sensitivity and specificity of sagittal STIR, PSIR, and FSE T2 in the detection of MS spinal cord lesions at 3T, in both the cervical and thoracic segments.

### MATERIALS AND METHODS

#### Subjects

Approval of the institutional review board to waive informed consent was obtained for this retrospective study. Inclusion criteria consisted of patients with definite or suspected MS who underwent MR imaging of the spinal cord at 3T, including sagittal STIR, PSIR, and FSE T2 as well as axial T2 sections in our institution from July 1 to October 31, 2013. Exclusion criteria were a final diagnosis of a disease causing spinal cord lesions different from MS and substantial motion artifacts. Spinal MR imaging of 39 consecutive patients was reviewed. One patient with a final diagnosis of Churg-Strauss syndrome was excluded. Thirty-eight patients were included in the study. In 11 patients, only the cervical segment was imaged, and in 12 patients, only the thoracic segment was covered. In 15 patients, both segments were included. This inclusion resulted in 26 cervical and 27 thoracic spinal segments available for evaluation.

#### MR Imaging Acquisition

Studies were performed on a 3T MR imaging scanner (Magnetom Skyra; Siemens, Erlangen, Germany) equipped with a 20-channel head-neck coil and a 16-channel spine-array coil. Image parameters are detailed in the Table. No contrast was administered before performing these sequences.

#### Image Analysis

Images of all subjects were stripped of patient identifiers and numbered randomly to facilitate blinded analysis. Two readers (P.A.-L. and L.A., neuroradiologists with 6 years’ experience in MR imaging) independently evaluated scans for the presence of spinal cord lesions on each spinal cord level on sagittal PSIR, sagittal STIR, and sagittal FSE T2 sequences. To facilitate this task and improve consistency between readers, we drew lines separating the spinal cord levels across the midpoint of the intervertebral disc, parallel to the vertebral endplates as shown in Fig 1C. These reference lines were available during all readings. The upper limit of the C1 segment was defined by the line connecting the tip of the odontoid and the upper edge of the posterior arch of C1. The lowest cord segment evaluated was T12, which was limited inferiorly by a line through the T12–L1 disc. Each sequence was evaluated independent of the other sequences so that the readers could not use them to confirm or rule out the presence of a lesion. Readings of different sequences for the same patient were separated by at least a 2-week interval to avoid memory bias. Disagreements were resolved by consensus. After the 2 independent readings were completed and the disagreements were resolved, an additional consensus reading was performed with all sequences available. This was considered the criterion standard. Combined information contained in sagittal FSE T2, proton density FSE, PSIR, and STIR and in axial gradient-echo T2 of the cervical spine and FSE T2 of the thoracic spine was taken into account to decide which spinal cord segments truly had demyelinating lesions and which were falsely identified as positive for demyelinating lesions due to the presence of artifacts. A lesion was considered a true one if visible on at least 2 sequences, regardless of the imaging plane.

In our experience, PSIR images of the thoracic spine often

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**Table: Acquisition parameters of MR imaging sequences**

| Segment | Sequence | Plane   | FOV (mm) | Acquisition Matrix | Thickness (mm) | TR (ms) | TE (ms) | TI (ms) | No. of Signals Acquired | Acquisition Time (min:sec) | Parallel Imaging |
|---------|----------|---------|----------|-------------------|----------------|---------|---------|---------|------------------------|-----------------------------|------------------|
| Cervical | PSIR     | Sagittal| 220      | 320 × 224         | 3              | 2400    | 9.4     | 400     | 2                      | 4:45                         | GRAPPA2         |
| Cervical | STIR     | Sagittal| 300      | 448 × 336         | 3              | 400     | 51      | 200     | 1                      | 5:46                         | GRAPPA2         |
| Cervical | FSE PD   | Sagittal| 220      | 320 × 288         | 3              | 2500    | 23      | 1       | 1                      | 2:02                         | None             |
| Cervical | FSE T2   | Sagittal| 220      | 384 × 307         | 3              | 3500    | 106     | 2       | 2                      | 3:35                         | None             |
| Cervical | GE T2    | Transverse| 180 | 256 × 218         | 3              | 740     | 17      | 2       | 2                      | 4:25                         | GRAPPA2         |
| Thoracic | PSIR     | Sagittal| 330      | 384 × 269         | 3              | 2400    | 9.5     | 400     | 2                      | 4:45                         | GRAPPA2         |
| Thoracic | STIR     | Sagittal| 330      | 448 × 336         | 3              | 4000    | 50      | 200     | 1                      | 5:46                         | GRAPPA2         |
| Thoracic | FSE PD   | Sagittal| 330      | 320 × 288         | 3              | 2500    | 21      | 1       | 1                      | 4:02                         | None             |
| Thoracic | FSE T2   | Sagittal| 330      | 448 × 336         | 3              | 4000    | 96      | 1       | 1                      | 4:26                         | None             |
| Thoracic | FSE T2   | Transverse| 180 | 256 × 176         | 5              | 6780    | 107     | 2       | 2                      | 4:46                         | None             |

Note: —GE indicates gradient-echo; PD, proton density; GRAPPA, generalized autocalibrating partially parallel acquisition.
have very poor quality, particularly in patients with a large thoracoabdominal diameter. To confirm our hypothesis that patient size affects PSIR sequence quality in the thoracic spine, we classified images by consensus into 2 groups: good quality or poor quality on the basis of noise and CSF signal saturation. We also measured the thickness of the dorsal back fat pad on PSIR sequences of the thoracic region. This was used as a surrogate marker of the thoracoabdominal diameter, which could not be measured directly on the images. This evaluation was performed by 1 author (P.A.-L.) in the midline at the level of T1–T2, T8–T9, and L1–L2. The mean of the 3 values was calculated. Thicknesses of the back fat pad of patients with good- and poor-quality PSIR of the thoracic spine were compared.

**Statistical Analysis**

Interrater reliability analysis by the Cohen κ coefficient was performed to determine segmental lesion detection consistency among raters. κ values were interpreted on the basis of the convention by Landis and Koch.20

Estimates of sensitivity and specificity were calculated for each sequence in the cervical and thoracic segments (On-line Table). To perform the calculation, we summated all false-positive, false-negative, true-positive, and true-negative segments of the different patients. Thus, calculated values of sensitivity and specificity do not represent directly the accuracy of the sequence in diagnosing MS but do represent the accuracy of the sequence in distinguishing an affected and nonaffected segment of the cord. Confidence intervals for sensitivity and specificity were produced with the Wilson score method.21 Accuracy of the sequences was also calculated.

Mean thickness of the thoracic back fat pad of the good-quality and poor-quality groups was compared by using the Mann-Whitney U test. The Pearson correlation coefficient was used to assess correlations between the thickness of the back fat pad and the sensitivity of the sagittal PSIR of the thoracic spine.

**RESULTS**

Twenty-six female and 12 male subjects were included (mean age, 42.1 years; range, 25–68 years). Of 38 subjects included in the study, 33 had clinically definite MS (n = 26, relapsing-remitting MS; n = 3, primary-progressive MS; n = 3, secondary-progressive MS; and n = 1, progressive-relapsing MS). The mean disease duration was 9 years (range, 1–27 years). The remaining 5 subjects included 2 patients with clinically isolated syndrome, 1 patient with radiologically isolated syndrome, and 2 patients with vague sensory symptoms without a definite clinically isolated syndrome.

There were no exclusions due to motion artifacts. As per criterion standard evaluation (consensus reading with all sequences available), spinal cord lesions were present in 84% of the cervical examinations and in 85% of the thoracic segments. In 6 patients, the upper thoracic region was not included in the FOV; thus, 8 thoracic segments could not be evaluated. One hundred eighty-two cervical and 316 thoracic spinal cord segments were evaluated. Of these, 106 cervical and 128 thoracic segments were affected by MS lesions. The number of true-positive, true-negative, false-positive, and false-negative spinal cord segments and the sensitivity, specificity, accuracy, and κ coefficient of each sagittal sequence are shown in the On-line Table. Comparisons among the number of true-positive, false-negative, and false-positive spinal cord segments of the 3 different sagittal sequences in the cervical and thoracic segments are shown in Fig 2. In the cervical region, the sensitivity of STIR and PSIR was significantly higher than that of FSE T2 (P < .05) (Fig 1). No significant difference was found in sensitivity between PSIR and STIR in the cervical region. In the thoracic region, the sensitivity of STIR was significantly higher than that of FSE T2 and PSIR, and the sensitivity of FSE T2 was significantly higher than that of PSIR (P < .05). Specificities of all cervical and thoracic sagittal sequences studied were very similar, with no significant differences identified. Interobserver agreement was higher for STIR than for FSE T2 and PSIR, in both the cervical and thoracic segments.

At visual inspection, all sequences were rated as good quality except for 10 PSIR thoracic studies that showed inhomogeneous CSF saturation and increased noise (Fig 3). The distribution of the variable “size of the back fat pad” in the good-quality and poor-quality groups is shown in Fig 3E. Back fat pad thickness in the poor-quality group was significantly higher compared with the high-quality group (P < .001). The mean back fat pad thickness was 13 mm (95% CI, 9.9–16.1 mm) in the good-quality group.
and 24.6 mm (95% CI, 21.8–27.3 mm) in the poor-quality group. A moderate negative correlation (Fig 3F) was found between the thickness of the back fat pad and the sensitivity of the sagittal PSIR sequence of the thoracic spine ($r = -0.58, P = .003$).

**DISCUSSION**

This study shows that lesion detection in the cervical region is lower on sagittal FSE T2 than on PSIR and STIR. The sensitivity of sagittal PSIR in our sample was slightly higher than that of STIR; however, the differences were not significant. In the thoracic spine, lesion detection of STIR was significantly better than that of FSE T2 and PSIR and lesion detection of FSE T2 was significantly better than that of PSIR.

The poor performance of PSIR in the thoracic region appears to be related to the thickness of the back fat pad. We are unsure about the cause of this effect in large patients. One explanation may be the dielectric effect. When the abdominal diameter exceeds the radiofrequency wavelength, the 180° pulse is not homogeneous through the FOV. Therefore, some areas may be getting an inversion pulse different from 180°. This effect, however, should also be detected on STIR because both are inversion recovery sequences. Another possibility may be a problem in phase unwrapping because the generation of true phase-sensitive images requires phase images to be
unwrapped first. The algorithm for phase unwrapping depends on a smooth variation in phase, and this constraint is sometimes not met, depending on the patient’s anatomy. A similar problem was found by Riederer et al when using a double inversion recovery sequence in the cervical cord. They excluded 10% of their sample due to artifacts presumably caused by adiposity. We did not observe this problem in the cervical examinations, but we found it in 37% of our thoracic examinations. We decided not to exclude large patients with poor image quality because our intention was to test sequences as they would be used in the routine clinical practice. Our observation is noteworthy because a sequence that provides suboptimal image quality in 37% of the patients would not be adequate for routine clinical use; thus, the utility of PSIR to detect thoracic spinal cord lesions may be limited.

Conventionally, sagittal FSE T2 is the most commonly used sequence when imaging the spinal cord in MS. Our results suggest that the use of an additional sagittal sequence can significantly increase the detection of MS lesions in the cervical and thoracic cord without a substantial decrease in specificity. Both sagittal PSIR and STIR perform well in this regard in the cervical spine, but only STIR appears to be adequate in the thoracic region. All the sequences studied have reasonable scanning times and can be implemented in routine clinical practice, but there is a small advantage for cervical PSIR because it is 1 minute shorter than STIR with our acquisition parameters.

We used a segment-based approach to quantify affected spinal cord tissue. With this method, interobserver agreement was moderate for PSIR and FSE T2 of the thoracic spine, substantial for FSE T2 and PSIR of the cervical spine and for STIR of the thoracic spine, and almost perfect for STIR of the cervical spine. Studies comparing lesion detection of different sequences in the spinal cord often had a consensus read or a single read or did not report values. Of note, our values using a segment-based approach to quantify spinal cord lesion load are higher than those reported in prior studies, which used lesion count; this finding suggests that a segment-based approach may be a more reliable way to quantify lesion burden in the spinal cord in MS.

The results of our study provide useful information for the design of 3T MR imaging protocols in MS. The superiority of STIR over T2 at 1.5T has been demonstrated on prior studies. Dietemann et al reported that the use of STIR in addition to sagittal T2 in the spinal cord increases the number of MS lesions detected by 46%. Another study by Nayak et al, which included a variety of scans performed at 1.5T and 3T, showed a 35% increase in the number of lesions detected using STIR compared with conventional sagittal T2. At 3T, Poonawalla et al reported improved lesion detection of sagittal STIR over sagittal FSE T2; however, the sample was small and no statistical analyses were performed to support the findings. In the study by Philpott and Brotchie, sagittal STIR showed a statistically significant increase in cervical cord lesion detection compared with FSE T2 at 3T. Our findings, therefore, are in agreement with previous studies at 1.5T and 3T that have shown that sagittal T2 alone does not have adequate sensitivity to detect cervical cord lesions (Fig 1), supporting the need to incorporate alternative sequences with higher sensitivity to accurately detect cervical cord lesions in MS. We have also demonstrated the utility of STIR sequences for improving spinal cord lesion detection in the thoracic segment at 3T, which has not been assessed in detail in prior studies.

Our findings support the use of PSIR as another alternative to improve the detection of MS lesions in the cervical spinal cord. In our study, PSIR was not superior to STIR in lesion detection in the cervical spine. Prior studies comparing these 2 sequences include the work by Poonawalla et al (cervical STIR versus PSIR) and Philpott and Brotchie (cervical T1 inversion recovery without phase reconstruction versus STIR); in both cases, samples were very small and statistical analysis of lesion detection was not performed. Thus, the existing literature does not provide enough evidence regarding the comparison between PSIR and STIR in the cervical cord, though a trend toward PSIR superiority has been observed, as in our study.

Our results raise the question of whether sagittal T2 should be included in the cervical examinations of patients with MS at 3T, given its inferiority compared with other sequences. We have shown the advantage of substituting sagittal T2 with STIR or PSIR in terms of cervical cord lesion detection; however, we have not evaluated the impact of eliminating sagittal T2 in the characterization of nondemyelinating cord pathologies and incidental findings.

This study has a number of limitations. First, we used a segment-based lesion-quantification approach in which the only information recorded was the presence or lack of lesions in each cord segment. Although this method showed substantially better interobserver agreement than lesion count, it has the disadvantage of being insensitive to differential lesion burden within a particular segment. Second, we focused on comparing FSE T2, STIR, and PSIR, but recent studies have demonstrated that double inversion recovery and T1 MPRAGE can be applied in the cervical cord of patients with MS; however, their long acquisition time (>7 minutes) limits their application in routine clinical practice. Due to the retrospective nature of our study, we did not have these sequences available for analysis in our study population, but future studies comparing T1-weighted phase-sensitive inversion recovery and T1 MPRAGE with STIR and PSIR would be useful. Third, contrary to prior studies, we did not calculate the body mass index. This parameter is easy to calculate and would allow rapid pre-scanning decision-making for STIR versus PSIR.

CONCLUSIONS
Our results suggest that at 3T, an additional sequence other than sagittal FSE T2 should be incorporated to improve lesion detection in the spinal cord of patients with MS. In the cervical cord, both sagittal STIR and PSIR showed significantly better lesion detection than sagittal FSE T2. In contrast, in the thoracic spinal cord, the sensitivity of PSIR is markedly worse than that of STIR and FSE T2, particularly in large patients, and STIR is clearly superior to FSE T2. These findings should be taken into account when planning sequence protocols to optimize lesion detection in the spinal cord in MS.
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