Chemical shift MR imaging in the lumbar vertebra: the effect of field strength, scanner vendors and flip angles in repeatability of signal intensity index measurement

Zebin Xiao†, Jian Li†, Chengqi Li‡, Yuyang Zhang†, Dejun She† and Dairong Cao†*

Abstract

Background: To evaluate the reproducibility of signal intensity index (SII) measurements with MRI systems from different vendors and with different field strengths, and to test the effectiveness of flip angle.

Methods: Thirty-two healthy volunteers (mean age 35.3 ± 9.3 years) were enrolled in this ethics committee-approved study. Chemical shift MR imaging was performed on 1.5- and 3.0-T MR systems from three vendors. Two independent observers measured SII values in five lumbar segments. Inter- and intraobserver agreement was assessed using the interclass correlation coefficients (ICCs). Differences of mean SII values between different field strengths and MR vendors as well as flip angles were compared by using repeated-measures analysis of variance. Differences of mean SII values between different flip angles were also compared by using paired-sample t test.

Results: Inter- and intra-observer correlation coefficients showed good agreement (all ICC > 0.75) when measuring SII values at different MR systems (ICCs ranging from 0.896 to 0.983) and flip angles (ICCs ranging from 0.824 to 0.983). There were no significant differences in mean SII values measured by different MR vendors with different field strengths (all p > 0.05 ranging from 0.337 to 0.824). The differences in the mean SII between the four different flip angles were statistically significant (all p < 0.05 ranging from < 0.001 to 0.004) except the group of flip angle 50° versus 70° (p = 0.116).

Conclusion: The SII measurement using chemical shift MR imaging may be comparable between different MR systems. Also high flip angles showed better stability to quantitate lumbar fat content.

Keywords: Chemical shift MRI, Repeatability, Bone marrow, In-phase and out-of-phase, Signal intensity index (SII)

Background

Chemical shift magnetic resonance (MR) imaging (also known as in-phase and out-of-phase imaging or opposed-phased imaging) is a simple technique that takes advantage of the fact that water and lipid hydrogen protons in a single voxel show slightly different precession frequencies [1]. Based on phase differences in images acquired via different TEs, lipid and water signals are additive on in-phase images and subtracted on opposed-phase images [2]. This technique has been proved to be extremely useful for characterization of lesions and organs with fatty components and has gained widespread acceptance [3–6]. In clinical practice, it is widely used to diagnose lipid-poor adrenal adenomas [3, 7]. On the same bias, chemical shift MR imaging has been used to evaluate vertebral bone marrow fat content in osteoporosis or in distinguishing benign and malignant causes of vertebral bone marrow infiltration [2, 4, 7–16]. Furthermore, some
investigators measured the signal intensity index (SII) value to avoid the problem of signal intensity variability produced by the reference tissue, and found that it appeared to be the most reliable method for differentiating adenomas from non-adenomas [7, 17]. However, there exists a major difference if measurements are performed at the adrenal gland or the bone marrow as reproducibility errors always have to be considered in relation to the variance that is expected in a population.

In spite of great interest in and enthusiasm about chemical shift MR imaging, there is a clear need for standardization of both the acquisition and the interpretation of chemical shift MR images to resolve current difficulties in comparing SII values from different studies or from different sites to enable validation of this quantitative parameter as a qualified biomarker in the context of multicenter studies.

SII measurements can be influenced by many factors such as chemical shift effect, susceptibility effect (i.e. T2* decay) and T1 relaxation, etc. [8, 18, 19]. Chemical shift effect occurs due to the slightly different precession frequency of water and fat. Susceptibility effect results from field inhomogeneity which is obvious at 3.0 Tesla or from the composition of the lesion itself [19]. As for the T1 relaxation, it depends on imaging parameters such as repetition time (TR) and flip angle [20]. Overall, whether SII values measured by chemical shift MR imaging can be compared across MR systems from different vendors and across field strengths remains an open question.

Thus, the aims of this study were to evaluate the reproducibility of SII values in the lumbar segments measured with MRI systems from different vendors and at different field strengths, and to test the effect of flip angle on assessing bone marrow fat content with respect to measurement of the signal intensity index (SII).

### Methods

This clinical study was approved by our Institutional Review Board (First Affiliated Hospital of Fujian Medical University) and written informed consent was obtained from all healthy volunteers enrolled in the study.

### Study population

In this present study, 35 healthy volunteers were included for chemical shift MR imaging of the lumbar spine between March 2015 and May 2015. The inclusion criteria for this study were as follows: (1) no history of trauma or surgery in the lumbar spine; (2) no history of acute or chronic pain in lower back; (3) no history of diseases which could change the signal intensity of lumbar marrow. The exclusion criteria was performed after interviewing the volunteers and reviewing their medical records, which included: (a) a history of or findings related to marrow diseases such as osteoporotic/traumatic fracture, traumatic, myeloma, osteosarcoma, lymphoma, spondylitis, etc.; (b) contraindications to MR imaging; (c) failure to complete the chemical shift imaging procedure for any reason; and (d) poor image quality insufficient for image analysis.

### MR imaging protocol

Data were acquired with 1.5 T MR systems (Vantage Alta, Toshiba Medical Systems, Otawara-shi, Japan) using phased-array spine coils with the following sequences: sagittal T1-weighted spin-echo sequence (500/10 [repetition time msec/echo time msec]), sagittal T2-weighted fast spin-echo sequence (4000/110 [repetition time msec/echo time msec]) and sagittal short inversion time inversion-recovery (STIR) fast spin-echo sequence (3500/65 [repetition time msec/echo time msec]). Chemical shift imaging data were acquired with 1.5 T MR

### Table 1 Intra- and interobserver agreement for SII measurement with different field strengths and vendors (Flip angle = 70°)

| Agreement | Siemens | GE | Toshiba |
|-----------|---------|----|---------|
| Reader 1/First time Reader 2/Second time | 0.935 (0.912, 0.952) | 0.961 (0.947, 0.973) | 0.972 (0.962, 0.989) |
| Intraobserver | 0.962 (0.945, 0.995) | 0.954 (0.983, 0.971) | 0.976 (0.967, 0.982) |
| Interobserver | 0.916 (0.885, 0.938) | 0.952 (0.935, 0.965) | 0.932 (0.878, 0.964) |

SII, Signal intensity index; Note. — Data in parentheses are 95% LOA

### Table 2 Intra- and interobserver agreement for SII measurement with different flip angles

| Agreement | Flip angle = 10° | Flip angle = 30° | Flip angle = 50° | Flip angle = 70° |
|-----------|-----------------|-----------------|-----------------|-----------------|
| Reader 1/First time Reader 2/Second time | 0.824 | 0.961 | 0.926 | 0.972 |
| Intraobserver | 0.668 (0.907) | 0.944 (0.983) | 0.854 (0.963) | 0.912 (0.952) |
| Interobserver | 0.858 | 0.969 | 0.923 | 0.954 (0.995) |
| flips | 0.881 | 0.913 | 0.936 | 0.896 |
| Observer | 0.796 (0.941) | 0.902 (0.941) | 0.879 (0.964) | 0.860 (0.923) |

Note. — Data in parentheses are 95% LOA
Fig. 1 (See legend on next page)
systems from two vendors (Vantage Atlas, Toshiba Medical Systems, Otawara-shi, Japan; Signa Twinspeed, GE Healthcare, Milwaukee, WI) and 3.0 T MR system from one vendor (Magnetom Verio, Siemens Medical Solutions, Erlangen, Germany) using phased-array spine coils in one week. The sequence parameters on 1.5 T MR systems were as follows: sagittal out-of-phase (OP) (2.4/192 [repetition time msec/echo time msec]) fast spoiled gradient echo MR imaging. And the sequence parameters on 3.0 T MR system were as follows: sagittal in-phase (IP) (2.4/192 [repetition time msec/echo time msec]) MR imaging. All chemical shift MR imaging data were transferred to an independent workstation for evaluation. Two readers (Z.X. and Y.Z., radiologists with 2 and 4 years of clinical experience in musculoskeletal MR imaging, respectively) independently drew rectangular regions of interest (ROIs) with a specific size in the center of three representative sections of each vertebral body, including midsagittal sections and adjacent two sections. The ROI size was initially defined by two authors in consensus (D.C. and J.L., radiologists with 28 and 23 years of clinical experience in musculoskeletal MR imaging, respectively) in a test set of images. The ROI was defined based on the largest possible rectangular size measuring 1.0 cm × 1.5 cm for the respective sections, and size for each region was kept constant during placement of ROIs by using the function of copy and paste in our workstation. Thus, a total of 54 ROIs were collected for each volunteer (three sections per vertebral body, five vertebral body, three vendors with two field strengths and MR vendors as well as flip angles were also compared by using paired-samples analysis). The ROI size was considered significant when p values were less than 0.05.

Results
Population demographics
Thirty-five volunteers were enrolled in this study. Three of them did not complete the study because of poor image quality (n = 2) or incomplete acquisition of all...
sequences (n = 1). Thirty-two volunteers successfully completed the imaging examinations (22 men and 10 women; mean age 35.3 ± 9.3 years; age range 21–50 years), and all the MR data sets were eligible for evaluation.

Repeatability of SII measurements in different MR systems and flip angles

The SIIIs measured in this study met the normal distribution (p = 0.193) and homogeneity of variance (p = 0.128). The intraobserver ICC calculated based on reader 1’s two measurements of SII values in 3.0 T Siemens system ranged from 0.896 to 0.972, with the 95% LOA ranging from (0.860, 0.923) to (0.962, 0.989) (Table 1). Interobserver agreement between reader 1’s first measurements and reader 2’s measurements was good for the three MR systems, with ICC (95% LOA) ranging from 0.916 (0.885, 0.938) to 0.983 (0.954, 0.975). The intra- and interobserver ICC results show good agreement between MR systems (all ICC > 0.75).

As shown in Table 2, the intra- and inter-observer ICC (95% LOA) ranged from 0.824 (0.668, 0.907) to 0.983 (0.954, 0.995), showing good agreement in measuring SIIIs in different flip angles (all ICC > 0.75).

Mutual agreement of mean SII measurements with different MR systems

The goal of this study was to assess agreement of mean SII values with different vendors at the same field strength (1.5 T Toshiba Vantage–1.5 T GE Signa), different field strengths and vendors (1.5 T GE Signa–3.0 T Siemens Verio; 1.5 T Toshiba Vantage–3.0 T Siemens Verio).

Overall, the agreement of above three conditions was desirable as the mean differences were very small (the mean differences ranged from −1.324 to 3.462) and most of the data points lied within 95% LOA (Figs. 1, 2). As shown in Table 3, the bias was not systematic, but depended on the specific lumbar segments and on the different MR systems. For instance, there existed significant bias on the measurements of SII values in L1 with three different MR systems; some segments (for example L3) showed negligible bias in comparison within the same field strength (Toshiba-GE), but a little bias when combining 1.5 T and 3.0 T data (1.5 T GE–3.0 T Siemens and 1.5 T Toshiba–3.0 T Siemens).
Comparison of mean SII values with three vendors in Two field strengths

The mean SII values of all lumbar vertebrae at 3.0 T Siemens, 1.5 T GE and 1.5 T Toshiba were 70.9 ± 5.4%, 71.3 ± 5.4%, and 71.5 ± 8.2%, respectively. Table 4 and Fig. 3 summarized the comparison of mean SII values with 3 MR systems in different lumbar segments. From the aspect of different field strength of MR systems, there were no significant differences in mean SII values (3.0 T Siemens vs 1.5 T GE, p = 0.337; 3.0 T Siemens vs 1.5 T Toshiba, p = 0.561). From the aspect of the same field strength of different vendors (1.5 T GE vs 1.5 T Toshiba), there were no significant differences in mean SII values (p = 0.824). Moreover, the mean SII values in the same lumbar segment had no significant differences among the measurement in the three different MR vendors (all p > 0.05, range from 0.26 to 0.95), and the mean SII values for each MR vendor at different lumbar segments had no significant differences either (all p > 0.05, ranging from 0.17 to 0.88).

Comparison of mean SII values in different flip angles

The mean SII of lumbar vertebra at 3.0 T Siemens with different flip angles (flip angle = 10, 30, 50, 70°) was 63.7 ± 7.4%, 67.7 ± 7.5%, 69.7 ± 6.0% and 71.0 ± 5.5%, respectively. The mean SII values showed a tendency of increasing with flip angles, and there were significant differences between different flip angles (p < 0.001) although there was no significant difference between the groups of flip angle 50° and 70° (p = 0.116) (Fig. 4, 5).

Table 3 Reproducibility of Mean SII Measurements in Each of the Five Lumbar Segments with Three Vendors in Two Field Strengths

| Lumbar Segment | 1.5 T Toshiba- 1.5 T GE | 1.5 T GE- 3.0 T Siemens | 1.5 T Toshiba - 3.0 T Siemens |
|---------------|------------------------|-------------------------|-------------------------------|
| L1            | 1.002 (−9.384, 11.388) | 2.460 (−18.067, 22.987) | 3.462 (−12.770, 19.693)       |
| L2            | −0.906 (−12.100, 10.289)| −0.327 (−18.893, 18.239) | −1.233 (−17.012, 14.547)     |
| L3            | −0.094 (−12.940, 12.751)| −1.230 (−16.663, 14.203) | −1.324 (−16.319, 13.671)     |
| L4            | −0.303 (−11.095, 10.488)| −0.327 (−14.208, 13.554) | −0.631 (−14.787, 13.526)     |
| L5            | −0.452 (−8.683, 7.779)  | 0.456 (−14.329, 15.242)  | 0.004 (−15.138, 15.142)      |

Note.—Data in parentheses are 95% LOA

Table 4 Comparison of Mean SII Measurement in Each of the Five Lumbar Segments with Three Vendors in Two Field Strengths

|                  | 3.0 T Siemens | 1.5 T GE | 1.5 T Toshiba | P Value |
|------------------|--------------|---------|--------------|---------|
| L1               | 70.124 ± 6.865 | 72.564 ± 10.121 | 73.505 ± 8.429 | 0.257   |
| L2               | 70.711 ± 5.721 | 71.156 ± 9.278  | 71.987 ± 8.255  | 0.810   |
| L3               | 71.223 ± 4.939 | 70.010 ± 8.401  | 69.917 ± 8.642  | 0.737   |
| L4               | 71.344 ± 4.953 | 71.077 ± 6.925  | 71.015 ± 6.909  | 0.930   |
| L5               | 71.225 ± 4.458 | 71.601 ± 6.882  | 71.658 ± 6.822  | 0.949   |

P Value* 0.882 0.171 0.643

Note.—Data are mean SIs (%) ± standard deviation

* Comparison of Mean SIs in the left- right direction

* Comparison of Mean SIs in the superior- inferior direction

Discussion

Chemical shift imaging is an useful technology in abdominal imaging by detecting lipid content [1, 3, 22, 26]. It was initially introduced for the assessment of bone marrow in 1985 by Wismer et al. [27], and also has been widely used in musculoskeletal imaging by measuring SII value for evaluation of bone marrow fat content or differentiation benign from malignant lesions [8, 9, 11–14]. But the lack of standardization in data analysis is a major challenge to the widespread and uniform use of chemical shift MR imaging in musculoskeletal imaging when comparing results of different studies. Furthermore, a reliable interpretation of the results contributed by different centers requires comparability of data acquired with MR systems at different institutions, which are often from different vendors and are operated at different field strengths. To the best of our knowledge, there are no previous reports describing the repeatability of SII measurements in different vendors, field strengths and flip angles.

In the present study, inter- and intraobserver correlation coefficients were good when measuring SII values with different MR systems (all ICC > 0.75), which indicates an excellent repeatability. Further study demonstrated that the mutual agreement of three MR systems was satisfying as the mean differences were very small and most of the data points lied within 95% LOA. Considering that imaging parameters are typically optimized for signal-to-noise ratio (SNR) and/or scan time, a given protocol may induce significant T1-weighting bias in the fat fraction estimate [18]. Our study has also shown that the bias was not systematic but depended on specific lumbar segments and on the different MR systems.

The mean SII values measured in our study for different lumbar segments lied within the previously reported range for all three vendors and for both field strength [15], but the mean SII values were slightly higher than the literature reported, which probably resulted from the effects of T1-weighting amplification induced by the high flip angle in our study. For the same field strength of different MR vendors, we did not find a significant difference in mean SII values in any of the evaluated lumbar segments. Furthermore, the agreement of mean SII values in the same field strength was better than different field strengths. All of these pointed to the
conclusion that quantitative analysis for lumbar fat content with chemical shift imaging in different MR vendors of the same field strength is comparable.

With the improvements in MR technology, the theoretical advantage of an increased SNR provided by the higher field strength is paralleled by disadvantages and challenges [19, 22]. Moreover, there exists fundamental differences in the MR physics of 3.0- and 1.5-T MR systems. As a result, chemical shift MR imaging at 1.5 T cannot be applied to 3.0 T MR imaging [19]. Thus, the repeatability of different field strengths is necessary to be researched. In this context, we found no significant difference in mean SII values with 1.5- and 3.0-T MR systems, and the repeatability of mean SII values in different field strength were good. Although the literature had reported that there were two factors influencing SI loss on OP images: chemical shift effect and susceptibility effect (i.e. T2* decay), and susceptibility effect occurred due to field strength inhomogeneity which was stronger at 3.0 T than at lower field strength [18, 19, 28], the results in our study indicated that quantitative analysis for lumbar fat content with chemical shift imaging in different field strengths MR systems may be comparable. Sebastian et al. had investigated the SII at 3.0- and 1.5-T MR imaging for prospectively quantitative analysis in a phantom study, and the result was similar to what we obtained.

In theory, the SI loss on OP images should be sensitive to differences in T1 relaxation, except for chemical shift effect and susceptibility effect [19]. With a poor choice of TE, susceptibility artifact on an OP image acquired later than an IP images can occur and may lead to the misinterpretation of a malignant adrenal lesion as a...
benign adenoma [22]. As a result, there were several studies focusing on effect of echo time in chemical shift MR imaging, but little or none literatures discuss the effects of T1 relaxation, which introduces a dependence on imaging parameters such as flip angle. In our current study, we did find no significant difference between the group of flip angle 50° versus 70° while there were significant differences between other groups. Moreover, the mean SII values increased with the augment of flip angle. The reason may be that a low flip angle could decrease the SNR leading to a bias in measuring SII and smaller estimates, and/or that due to the shorter T1 of fat, a low flip angle would influence full T1 recovery and make the ratio TE/TR changed.

The result of our study indicated that chemical shift MR imaging might be applicable as a biomarker in the lumbar spine, even in multicenter studies combining different vendors and different field strengths. For instance, studies that focus on treatment-induced SII value differences can be performed with any combination of MR systems. Ongoing research could focus on providing correction factors for intervendor or interfield strength comparisons or further optimizations of SNR to reduce overall test-retest variability.

Our study still have some limitations. Firstly, we included healthy volunteers instead of a phantom for measurements. But we aimed to simulate a clinical environment similar to clinical practice for our analysis. Secondly, we did not evaluate the reproducibility in subjects with pathology to make meaningful comparisons. A large number of patients with different diseases in various vertebrae or organs would be necessary. However, it may be difficult to acquire a large number of patients with the disease who would be willing to undergo three repeated measurements. Thirdly, subjected to the limited conditions, we did not strictly compare mean SII values...
between different filed strength at the same vendor. A next multicenter study will be performed to refine our research. Lastly, as SIs measured in chemical shift MR imaging are not quantitative, quantitative MR measurement including T1 and T2 relaxation times, such as a proton density fat fraction (PDFF) should be intended in our further research.

Conclusion
In conclusion, the signal index (SII) using chemical shift MR imaging may be comparable between MR systems from different vendors and at different field strengths. In addition, high flip angles (50° or 70°) showed better stability for quantitative analysis of lumbar fat content, indicating that high flip angles should be chosen when other parameters are fixed.

Abbreviations
FA: Flip angle; ICC: Interclass correlation coefficients; IP: In phase; LOA: limits of agreement; MRI: Magnetic resonance imaging; OP: Out of phase; PDFF: Proton density fat fraction; SII: Signal intensity index; SNR: Signal-to-noise ratio; STIR: Short inversion time inversion-recovery; TR: Repetition time

Acknowledgements
None.

Funding
None.

Availability of data and materials
Due to statutory provisions regarding data- and privacy protection, the dataset supporting the conclusions of this article is available upon individual request directed to the corresponding author.

Authors’ contributions
ZX and JL participated in the design of the study, performed the experiments and the statistical analysis and drafted the manuscript. DC participated in the design of the study, performed the experiments and the statistical analysis. YZ participated in the design of the study and assisted with manuscript preparation. CL participated in the design of the study and assisted with manuscript preparation. DS assisted with manuscript preparation. All authors read and approved the final manuscript.

Competing interests
The authors declare that they have no competing interests.

Consent for publication
Not applicable.

Ethics approval and consent to participate
The protocols were reviewed and approved by the Ethics Committee of our University Hospital and was conducted in accordance with the Declaration of Helsinki. All patients gave written informed consent to participate in this study.

Author details
1Department of Radiology, First Affiliated Hospital of Fujian Medical University, 20 Cha-Zhong Road, Fuzhou, Fujian 350005, China. 2Department of Radiology, Samning Hospital of Integrated Traditional and Western, Sanming, Fujian 365000, China.

Received: 27 May 2016 Accepted: 14 November 2016 Published online: 25 November 2016

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