SHORT REPORT

CT-like images of the sacroiliac joint generated from MRI using susceptibility-weighted imaging (SWI) in patients with axial spondyloarthritis

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ABSTRACT

BACKGROUND To analyse the added value of susceptibility-weighted imaging (SWI) compared with standard T1-weighted (T1) MRI for detecting structural lesions of the sacroiliac joint (SIJ) in patients with axial spondyloarthropathy (axSpA) using CT as reference standard.

MATERIAL AND METHODS Sixty-eight patients with suspected or proven axSpA underwent both MRI and CT of the SIJ on the same day. Two readers separately scored CT, T1 and SWI for the presence of erosions, sclerosis and joint space changes using an established 24-region SIJ model. Disagreement was resolved by a third reader. Diagnostic accuracy (McNemar test), Cohen’s kappa (κ), sensitivity (SE) and specificity were calculated on the joint level using CT as reference.

RESULTS In CT, 38 joints showed erosions, 67 sclerosis and 37 joint space changes. Agreement with CT for erosions was 92.6% (κ=0.811 (0.7–0.92)) in SWI and 87.5% (κ=0.682 (0.54–0.82)) in T1 (p=0.143) and agreement for sclerosis 84.6% (κ=0.69 (0.57–0.81)) and 62.5% (κ=0.241 (0.13–0.35)) (p<0.001), respectively. This resulted in superior SE of SWI (81.6% vs 73.7%) for erosions and sclerosis (74.6% vs 23.9%) at a minor expense of SP. No differences were detected for joint space changes.

CONCLUSION In patients with axSpA, SWI depicts erosions and sclerosis more accurately than T1 spin echo MRI at 1.5 T.

INTRODUCTION

Detection of structural damage of the sacroiliac joint (SIJ) including erosions, sclerosis and joint space changes is crucial for establishing the diagnosis of axial spondyloarthritis (axSpA) and monitoring disease progression. The modified New York Criteria include these structural changes in radiography but do not incorporate MRI.1 Conversely, the Assessment of SpondyloArthritis international Society criteria incorporate osteitis in MRI for classification but attach less weight to structural lesions.1–3 While conventional sequences have recently been shown to detect erosions with high diagnostic accuracy,4 standard MRI is limited because it cannot depict cortical bone directly (it is hypointense in all sequences) but relies on the contrast of bone, bone marrow and cartilage. Conversely, radiography and CT depict the bone directly due to X-ray attenuation of bone but involve radiation exposure and do not capture active inflammation, which is deemed important for early diagnosis but less specific for differential diagnosis.

There are some developments in MRI for direct bone depiction. One of them, susceptibility-weighted imaging (SWI),5 has been successfully applied to erosion detection in patients with peripheral arthritis.6 SWI depicts calcium structures directly by detecting and quantifying small magnetic field inhomogeneities surrounding calcium atoms. While SWI has been standard in many brain MRI protocols for several years, its transfer to musculoskeletal imaging was rather recent.7 Inversion of these images creates the impression of CT images in MRI.

The aim of our study was to investigate SWI in detecting structural SIJ damage in patients...
with axSpA compared with T1-weighted (T1) sequences and using CT as standard of reference.

### METHODS

#### Patients

We prospectively included 75 patients with suspected or diagnosed axSpA examined between February 2018 and November 2019. They were referred by the local rheumatology department either to confirm axSpA in patients with inflammatory back pain and other typical features of axSpA or to evaluate inflammatory activity in patients with an established diagnosis. The final diagnosis was made by the rheumatologist. Exclusion criteria were contraindications to MRI (eg, pacemaker), claustrophobia and pregnancy.

#### Imaging

All patients underwent MRI at 1.5 T. The protocol included conventional T1-weighted (T1) and STIR sequences and an SWI sequence of the SIJ acquired with 4 mm slice thickness and 10% gap between slices in oblique coronal orientation (online supplemental file 1). CT was performed on the same day as dual-energy CT (DECT) on a single-source scanner (Canon Aquilion One Vision) and served as standard of reference. DECT source data were reconstructed in 120-equivalent blended images (equivalent to conventional CT) and reformatted to 3 mm oblique coronal image stacks.

#### Image reading

Standard MRI (T1, STIR), SWI and CT of the SIJ were anonymised separately using Horos (The Horos Project, V.3.3.6, Pureview, Maryland, USA). Inverted SWI was viewed to imitate the impression of CT. The images were scored separately by two readers (reader 1: musculoskeletal radiologist with 11 years of experience; reader 2: research student with 2 years of experience). Disagreement was solved by an expert adjudicator (reader 3: musculoskeletal radiologist with 21 years of experience). Images were scored using a 24-region approach for erosions and sclerosis previously established by our group. Briefly, this model divides each SIJ into four quadrants and each quadrant into an anterior, central and posterior region. Joint space changes were scored per joint and region. Scores were as follows: 0–3 for erosions (0 no erosions, 1 small isolated erosions (n=1–2), 2 definite erosions (n=3–5; <3 mm), 3 multiple (n>5) or confluent erosions); 0–2 for sclerosis (0 no sclerosis, 1 minor sclerosis (5–9 mm), 2 evident sclerosis (≥10 mm)); and 0–4 for joint space changes (0 normal joint space, 1 possible widening/narrowing, 2 definite widening/narrowing, 3 partial ankylosis, 4 complete ankylosis). Scoring results were reported using an in-house developed online electronic Case Report Form (eCRF). Readers 1 and 2 scored 10 standard MRI and CT test cases before the study reading.

#### Data analysis

Erosions, sclerosis and joint space changes were defined as positive if a score of 2 or higher was assigned per joint by both readers. Any disagreement was resolved by the third reader. Sum scores were separately calculated for each joint and reader. The mean of both readers’ sum scores was calculated for each lesion in T1 and SWI for comparison with CT using Pearson’s r. Sensitivity (SE), specificity (SP) and likelihood ratios (LR+ and LR−) were calculated for each MRI protocol and each lesion. Diagnostic accuracy was calculated for each lesion in SWI and T1 using CT as reference, and the results were compared with a McNemar test. Cohen’s kappa (k) was calculated for agreement with CT and for interrater reliability. Statistical analysis was performed using SPSS (V.27.0.0.0).

### RESULTS

#### Patients

Seven of 75 patients were excluded from analysis because they did not undergo MRI (known claustrophobia n=2, cancelled MRI due to claustrophobia n=2, cancelled MRI due to back pain n=1, possible pregnancy n=1) or CT (technical error n=1). Further patient characteristics are presented in table 1.

#### Image reading and data analysis

Contingency tables, SE (SE), SP, LR and diagnostic accuracy data for the three types of structural lesions are compiled in table 2. Imaging examples are presented in figure 1.

#### Erosions

Thirty-eight joints were considered positive for erosions in CT, 35 in T1 and 34 in SWI. The sum score (average of both readers) was 2.05±3.37 in CT, 1.48±2.0 in T1 and 2.37±3.39 in SWI. Correlation with CT was moderate for T1 (r=0.786) and very strong for SWI (r=0.87). Diagnostic accuracy results including Cohen’s kappa for agreement with CT are presented in table 2. Inter-rater reliability was substantial for CT (k=0.741 (0.61–0.87); p<0.001), slight for T1 (k=0.185 (0–0.38); p<0.001) and moderate for SWI (k=0.424 (0.3–0.62); p<0.001).

#### Sclerosis

For sclerosis, 67 joints were scored positive in CT, 16 in T1 and 55 in SWI. The mean sum score for sclerosis was 2.87±3.09 in CT, 0.88±1.62 in T1 and 3.9±4.36 in SWI. Correlation with CT was moderate for T1 (r=0.677) and very strong for SWI (r=0.87). For diagnostic accuracy see table 2. Inter-rater reliability for the presence of sclerosis was moderate for CT (k=0.569 (0.44–0.7); p<0.001), substantial for T1 (k=0.701 (0.51–0.89); p<0.001) and fair for SWI (k=0.394 (0.25–0.55); p<0.001).

#### Joint space changes

Joint space changes were considered to be present in 37 joints in CT, 33 in T1 and 38 in SWI. The mean sum score for this change was 2.26±3.26 in CT, 1.58±2.65 in
T1 and 2.03±2.71 in SWI. Correlation of MRI with CT for joint space changes was very strong with both protocols (r=0.832 for T1, r=0.814 for SWI). For diagnostic accuracy see table 2. Cohen's kappa for inter-rater reliability was substantial for CT (k=0.669 (0.54–0.8); p<0.001) and T1 (k=0.654 (0.5–0.8); p<0.001) and fair for SWI (k=0.327 (0.16–0.5); p<0.001).

**DISCUSSION**

This is the first study using SWI to create CT-like MR images for detection of structural SIJ lesions in axSpA. Our results show that SWI improves the detection of erosions and sclerosis as important structural lesions of the SIJ, in terms of both diagnostic accuracy and

### Table 1  Patient characteristics

<table>
<thead>
<tr>
<th></th>
<th>All patients</th>
<th>Patients with inflammatory disease (n=40)</th>
<th>Patients with non-inflammatory disease (n=28)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Sex</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sex</td>
<td>41.2% female (28/68)</td>
<td>27.5% female (11/40)</td>
<td>60.71% female (17/28)</td>
</tr>
<tr>
<td><strong>Age</strong></td>
<td>40.54±12.23</td>
<td>39.85 (SD 12.23)</td>
<td>41.54 (SD 11.96)</td>
</tr>
<tr>
<td>HLA-B27 positivity</td>
<td>55.8% (29/52)</td>
<td>73.33% (22/30)</td>
<td>21.82% (7/22)</td>
</tr>
<tr>
<td>Mean CRP</td>
<td>12.34 (n=24)</td>
<td>14.89 (SD 17.78; n=15)</td>
<td>7.58 (SD 18.38; n=9)+negative in 3 patients</td>
</tr>
<tr>
<td><strong>BASDAI</strong></td>
<td>4.67±1.61 (n=33)</td>
<td>4.6 (SD 1.61; n=27)</td>
<td>5.02 (SD 1.61; n=6)</td>
</tr>
<tr>
<td>Modified New York Criteria positivity</td>
<td>N/A</td>
<td>52.5% (21/40)</td>
<td>N/A</td>
</tr>
<tr>
<td>ASAS MRI criteria positivity</td>
<td>N/A</td>
<td>52.5% (21/40)</td>
<td>N/A</td>
</tr>
<tr>
<td>Erosion sum score</td>
<td>2.05 (SD 3.37)</td>
<td>3.16 (SD 3.37)</td>
<td>0.46 (SD 3.26)</td>
</tr>
<tr>
<td>Sclerosis sum score</td>
<td>2.87 (SD 3.09)</td>
<td>3.26 (SD 3.09)</td>
<td>2.31 (SD 3.03)</td>
</tr>
<tr>
<td>Joint space sum score</td>
<td>2.26 (SD 3.26)</td>
<td>3.53 (SD 3.26)</td>
<td>0.45 (SD 3.24)</td>
</tr>
</tbody>
</table>

Patient characteristics are presented for all patients and by subgroup according to the rheumatologist’s final diagnosis (inflammatory vs non-inflammatory disease) by the rheumatologist. Inflammatory conditions were axial spondyloarthritis (axSpA) (n=35; r-axSpA: n=29, nr-axSpA: n=6), psoriatic arthritis with axial inflammation (n=4) and SAPHO (n=1). Non-inflammatory conditions were degenerative spine disease (n=14), osteitis condensans ili (n=13) and psoriatic arthritis without axial involvement (n=1). The sum scores are mean scores in CT. ASAS, Assessment of SpondyloArthritis international Society; BASDAI, Bath Ankylosing Spondylitis Disease Activity Index; CRP, C-reactive protein; SAPHO, synovitis, acne, pustulosis, hyperostosis and synovitis syndrome.

### Table 2  Cross table, sensitivity (SE), specificity (SP), likelihood ratio (LR) and diagnostic accuracy

<table>
<thead>
<tr>
<th></th>
<th>Erosions</th>
<th>Sclerosis</th>
<th>Joint space changes</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CT+</td>
<td>CT−</td>
<td>Diagnostic accuracy</td>
</tr>
<tr>
<td><strong>Erosions</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T1+</td>
<td>28</td>
<td>7</td>
<td>SE 73.68%</td>
</tr>
<tr>
<td>T1−</td>
<td>10</td>
<td>91</td>
<td>SP 92.86%</td>
</tr>
<tr>
<td>SWI+</td>
<td>31</td>
<td>3</td>
<td>SE 81.58%</td>
</tr>
<tr>
<td>SWI−</td>
<td>7</td>
<td>95</td>
<td>SP 96.94%</td>
</tr>
<tr>
<td><strong>Sclerosis</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T1+</td>
<td>16</td>
<td>0</td>
<td>SE 23.88%</td>
</tr>
<tr>
<td>T1−</td>
<td>51</td>
<td>69</td>
<td>SP 100.0%</td>
</tr>
<tr>
<td>SWI+</td>
<td>50</td>
<td>5</td>
<td>SE 74.63%</td>
</tr>
<tr>
<td>SWI−</td>
<td>17</td>
<td>64</td>
<td>SP 92.75%</td>
</tr>
<tr>
<td><strong>Joint space changes</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T1+</td>
<td>26</td>
<td>7</td>
<td>SE 70.27%</td>
</tr>
<tr>
<td>T1−</td>
<td>11</td>
<td>92</td>
<td>SP 92.93%</td>
</tr>
<tr>
<td>SWI+</td>
<td>28</td>
<td>10</td>
<td>SE 75.68%</td>
</tr>
<tr>
<td>SWI−</td>
<td>9</td>
<td>89</td>
<td>SP 89.9%</td>
</tr>
</tbody>
</table>

All values were calculated using CT as standard of reference or in comparison with CT (absolute agreement and Cohen’s kappa). Agreement of SWI with CT was significantly higher for sclerosis (p<0.001) and tended to be higher for erosions (p=0.143), while there was no difference for joint space changes (p=1).

Bold text have been used to mark the subsection headings of the table. SWI, susceptibility-weighted imaging.
correlation of sum scores. The detection of joint space alterations, however, is not improved by SWI compared with conventional MRI.

SWI has been shown to improve erosion depiction in patients with hand arthritis.\(^6\) While other novel MRI sequences such as volumetric interpolated breath-hold examination (VIBE) and MRI-based synthetic CT have been shown to be superior to T1 in detecting erosions in the SIJ,\(^9-11\) they do not allow direct visualisation of the cortical and trabecular bone structure and, thus, still suffer from typical MRI shortfalls. VIBE, for example, is a gradient echo sequence with an undesired T2* effect that causes a signal loss in the vicinity of calcium crystals, resulting in the typical stark contrast of soft tissue and bone. Conversely, SWI exploits the paramagnetic characteristics of calcium and the resulting T2* effect directly for improved contrast in musculoskeletal imaging. While SWI is a widely available and commonly used pulse sequence in neuroimaging and can be easily transferred to musculoskeletal imaging, MRI-based synthetic CT needs specifically trained AI software. Further developments in SWI have been used to directly quantify materials, but this was not investigated in our study.\(^12\)

SWI may have several advantages for clinical practise. Before the advent of SWI, evaluation of structural changes by direct depiction of cortical bone was only possible with imaging modalities using ionising radiation.\(^10\) A standard MRI protocol supplemented by SWI both provides diagnostic information on the presence of active bone marrow lesions (osteitis) and allows accurate detection of structural lesions in a single imaging session by adding 5 min and 47 s of scan time. This plays an important role when it comes to differential diagnosis or monitoring of disease progression.\(^14\) In MRI of the SIJ, an important differential diagnosis for the presence of osteitis is mechanical stress, for example, in osteitis condensans ilii. One factor that can help in differentiating these non-erosive conditions from axSpA is the absence of erosions in the presence of sclerosis.\(^15,16\) The latter is even more pronounced in SWI, but showed high specificity in our analysis.

Still, some limitations of this study have to be discussed. We included a mixed population of patients with and without axSpA. Ethical concerns prohibited inclusion of healthy controls due to the radiation exposure of CT. A statistically significant improvement was only shown...
for sclerosis. Our results need to be verified in larger patient populations, for MRI at 3 T, and in comparison with other novel sequences (VIBE, MR-based synthetic CT) and thinner slices. Some clinical and laboratory data were not available for all patients. We provide a structural lesion analysis only and do not elaborate on the diagnostic impact of SWI.

In conclusion, SWI depicts erosions and sclerosis more accurately than T1-weighted spin echo MRI at 1.5 T and may provide useful additional information for the diagnosis of axSpA.

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**Contributors** DD: patient acquisition, image scoring, data evaluation, statistical calculations, article draft, critical revision of the manuscript for important intellectual content. KGH: conception and design of the study, design of scoring system, image scoring, critical revision of the manuscript for important intellectual content. TD: patient acquisition, data collection, critical revision of the manuscript for important intellectual content. DP: patient acquisition, statistical calculations, critical revision of the manuscript for important intellectual content. MP: patient acquisition, critical revision of the manuscript for important intellectual content. MRM: conception and design of the study, critical revision of the manuscript for important intellectual content. TD: conception and design of the study, design of scoring system, image scoring, data evaluation, statistical calculations, critical revision of the manuscript for important intellectual content.

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