Fully automated joint space width measurement and digital X-ray radiogrammetry in early RA

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ABSTRACT

Objectives To study fully automated digital joint space width (JSW) and bone mineral density (BMD) in relation to a conventional radiographic scoring method in early rheumatoid arthritis (eRA).

Methods Radiographs scored by the modified Sharp van der Heijde score (SHS) in patients with eRA were acquired from the SWEdish FarmacOTherapy study. Fully automated JSW measurements of bilateral metacarpals 2, 3 and 4 were compared with the joint space narrowing (JSN) score in SHS. Multilevel mixed model statistics were applied to calculate the significance of the association between JSW and BMD over 1 year, and the JSW differences between damaged and undamaged joints as evaluated by the JSN.

Results Based on 576 joints of 96 patients with eRA, a significant reduction from baseline to 1 year was observed in the JSW from 1.69 (±0.19) mm to 1.66 (±0.19) mm (p<0.01), and BMD from 0.583 (±0.068) g/cm² to 0.566 (±0.074) g/cm² (p<0.01). A significant positive association was observed between ΔJSW and ΔBMD over 1 year (p=0.0001). On an individual joint level, JSWs of undamaged (JSN=0) joints were wider than damaged (JSN>0) joints: 1.68 mm (95% CI 1.70 to 1.67) vs 1.54 mm (95% CI 1.63 to 1.46). Similarly the unadjusted multilevel model showed significant differences in JSW between undamaged (1.68 mm (95% CI 1.72 to 1.64)) and damaged joints (1.63 mm (95% CI 1.68 to 1.58)) (p=0.0048). This difference remained significant in the adjusted model: 1.66 mm (95% CI 1.70 to 1.61) vs 1.62 mm (95% CI 1.68 to 1.56) (p=0.042).

Conclusions To measure the JSW with this fully automated digital tool may be useful as a quick and observer-independent application for evaluating cartilage damage in eRA.

Trial registration number NCT00764725.

INTRODUCTION

Despite technological advancements and the availability of ultrasound (US) and MRI modalities, conventional radiography (CR) remains the main imaging tool for rheumatoid arthritis (RA). The clinical use of radiographs as permanent medical records has several strengths, that is, the technology is accessible globally, they may be evaluated at any time, the severity of structural damage and progression can be assessed, and treatment effects determined. RA inflammation has a predilection for small joint involvement, and radiographs of the metacarpophalangeal (MCP) and metatarsophalangeal joints display radiographic progression particularly well. Joint space narrowing (JSN) due to destroyed cartilage may have a larger effect...
on functional status than erosions, making it a valid target for treatment.  

In RA clinical trials, the van der Heijde modification of the Sharp score (SHS) is currently the most frequently used scoring method for evaluating radiographic progression with JSN and erosions of selected joints in hands and feet. Although this scoring method has many strengths, its weaknesses are that the scoring is time-consuming and that it requires specialised training that seldom is available to the rheumatologists. In order to minimise the aforementioned limitations, semiautomated and fully automated software to better evaluate joint space width (JSW) have been developed and reviewed. The benefits of a fully automated program include higher reproducibility, faster results, observer independence and a higher sensitivity. Moreover, bone mineral density (BMD) measured with digital X-ray radiogrammetry (DXR) is a recent tool that quantifies BMD in metacarpals 2, 3 and 4. DXR has been proven sensitive in measuring BMD loss, and BMD loss has been shown to be associated with an increased risk of radiographic progression. Rapid loss of BMD is predictive of radiographic progression, and this has been shown already after 4 months. Using the SWEdish FarmacOTherapy (SWEFOT) study data of patients with early RA, we now investigate how JSW compares with JSN and explore the association between BMD and JSW using a multilevel mixed model, which has not previously been done. We also discuss the utility of these new tools in clinical practice.

METHODS  

Patients and study design  

Data of patients with early RA were acquired from the SWEFOT study. The detailed study design has previously been reported and is summarised in the online supplementary material. Of the 487 patients in the SWEFOT study, 119 patients met the inclusion criteria for this study (figure 1) and constitute the patients in the multilevel mixed model. Furthermore, 96 patients had both baseline and follow-up radiographs, which allowed for comparisons of progress over 12 months. A flow chart illustrating the selection process is shown in figure 1.

Radiographic assessment  

Radiographs of the hands and feet were obtained at baseline and 1 year.

Radiographic positioning of the hand  

Radiographic examinations were performed using a posteroanterior (PA) projection of the hand. The PA projection is the best conventional view for demonstrating malalignment, JSN and soft-tissue abnormalities in early RA. Bilateral hands are generally X-rayed, with the contralateral image used for bony structure comparison.

Technical factors  

An image receptor (IR), 10×12 inch (24×30 cm) crosswise for two or more images on one cassette, was used. A digital screen, lead masking and collimation is used for radiation protection and avoidance of scatter. A range of 50–60 kVp and 3–4 mAs were used, as well as a minimum source to image distance (SID) of 100 cm.

Positioning for PA projection  

First the patient is asked to sit at the end of the radiographic table that is adjusted to the patient’s height so that the forearm is resting on the table. Second, the patient’s forearm is well rested on the table with the hands placed with the palmar surface flat onto the cassette. Then the radiographic plate/cassette is centred to the MCP joints, and adjusted to the long axis of the cassette parallel with the long axis of the hand and forearm. Then the patient is asked to spread the fingers slightly to ensure correct positioning. The patient is then asked to relax the hand to avoid motion. Adhesive tape or positioning sponges can be used to prevent involuntary movement. A sandbag may be placed over the distal forearm if necessary. Finally,
a central ray is directed so that it is perpendicular to the third MCP joint.

**Evaluation criteria**

First, all fingers, the wrist and about 2.5 cm of the distal forearm should be visible. Second the MCP and interphalangeal joints (clear joint spaces) should appear open, indicating correct central ray location and that the hand was fully pronated. The long axis of the IR should be well aligned to the long axis of the hand and wrist. Third, free from positioning error, both sides of the hands should appear symmetrical unless pathology is suspected with no evidence of rotational error. The concavities of the shafts of the metacarpals and the phalanges 2–5, and the amount of soft tissue on either side of phalanges 2–5, should appear equal unless pathology is suspected. The digits should not overlap, and the fingers should be separated slightly with soft tissues clearly delineated.

These images of the hands, per standardised radiography protocol, are the ones used for radiographic assessment by both the clinician and the fully automated JSW program. Only images using the same modality type and settings at baseline and follow-up were analysed. Radiographic joint damage was assessed according to the SHS method, by visual inspection of the radiographs and using a semiquantitative scale of 0–4 for JSN. The radiographs were read in chronological order by one of two experienced readers blinded to clinical data. The interclass correlation coefficient was 0.94 and the smallest detectable difference (SDD) in SHS was 5.8 units.

**Fully automated JSW measurements**

Computer-assisted automated measurements of the MCP joint spaces were calculated from the digital hand radiographs of each patient using dedicated software (dxr-online, Sectra, Linköping, Sweden). This software was a further development of a semiautomated version used previously. The detailed method for measuring JSW is described in the online supplementary material. The measurement regions are illustrated in figure 2. The short-term in vivo reproducibility of JSW was tested with 30 healthy individuals using the same machine, radiographer and protocol at three separate time points on the same day. The hand was repositioned, that is, moved, between the three images. This was expressed as the coefficient of variation (CV%), which was 1.4% and the SDD was 0.062 mm for MCP 2, 3 and 4 of the 30 healthy volunteers.

**Metacarpal BMD**

In the DXR-BMD method, the narrowest part of the metacarpals 2, 3 and 4 is located. The cortical thickness and bone width are measured within the measurement regions. By assuming that bone density is constant and the bone elliptical, the BMD can be calculated in g/cm². The measurement regions are shown in figure 2.

**Statistics**

Normal distribution was tested using the Shapiro-Wilk test. For unpaired data, comparison between groups was performed using the unpaired t-test for normally distributed data, and the Mann-Whitney test for non-normally distributed data. For paired data, we used the paired t-test in case of normally distributed data, and the Wilcoxon test for non-normally distributed data, to determine whether the change over 12 months was significant. A multilevel mixed model was used to explore if there was a significant association between ΔJSW and ΔBMD over 1 year. This model clustered ΔJSW and ΔBMD on patient level and each hand separately. It also took into account repeated visits, that is, baseline and 1 year follow-up, and adjusted for height, age, gender and BMI. Additional multilevel mixed models were used to establish the difference in JSW between individual joints with cartilage damage (JSN score >0) versus joints with no cartilage damage (JSN score=0). These multilevel models took into account the correlations between the joints from the same hand as well as with the contralateral joints, since these joints are intrinsically related to each other. In these models, JSW was considered the dependent variable, and cartilage damage in the form of JSN the explanatory variable. We constructed models taking into account baseline and
Table 1  Demographic data of the whole SWEFOT population versus the study cohort

<table>
<thead>
<tr>
<th>Demographic</th>
<th>SWEFOT patients (n=487)</th>
<th>Study patients (n=119)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female sex, n (%)</td>
<td>344 (70.6)</td>
<td>93 (78.2)</td>
<td>0.13</td>
</tr>
<tr>
<td>ACPA pos, n (%)</td>
<td>310 (63.7)</td>
<td>60 (57.7)</td>
<td>0.25</td>
</tr>
<tr>
<td>RF pos, n (%)</td>
<td>333 (68.4)</td>
<td>76 (65.0)</td>
<td>0.47</td>
</tr>
<tr>
<td>DAS28, mean (SD)</td>
<td>5.73 (1.01)</td>
<td>5.64 (0.95)</td>
<td>0.39</td>
</tr>
<tr>
<td>ESR (mm), mean (SD)</td>
<td>39.8 (28.1)</td>
<td>37.5 (24.3)</td>
<td>0.45</td>
</tr>
<tr>
<td>CRP (mg/L), mean (SD)</td>
<td>33.7 (42.4)</td>
<td>34.8 (38.1)</td>
<td>0.69</td>
</tr>
<tr>
<td>HAQ, mean (SD)</td>
<td>1.19 (0.58)</td>
<td>1.18 (0.56)</td>
<td>0.95</td>
</tr>
<tr>
<td>SHS, mean (SD)</td>
<td>4.54 (8.01)</td>
<td>4.49 (8.20)</td>
<td>0.74</td>
</tr>
<tr>
<td>JSN, mean (SD)</td>
<td>2.63 (7.08)</td>
<td>2.45 (5.68)</td>
<td>0.50</td>
</tr>
<tr>
<td>Erosion score, mean (SD)</td>
<td>1.91 (3.75)</td>
<td>2.03 (3.70)</td>
<td>0.34</td>
</tr>
</tbody>
</table>

ACPA, anticitrullinated peptide antibody; CRP, C reactive protein; DAS28, Disease Activity Score of 28 joints; ESR, erythrocyte sedimentation rate; HAQ, Health Assessment Questionnaire; JSN, joint space narrowing; Pos, positive; RF, rheumatoid factor; SHS, modified Sharp van der Heijde Score; SWEFOT, SWEdish FarmacOTherapy.

Table 2  JSW, BMD, erosion score, JSN score and SHS at baseline and 12 months

<table>
<thead>
<tr>
<th>Measure (mean±SD)</th>
<th>Baseline (n=96)</th>
<th>12 Month follow-up (n=96)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Left hand BMD (g/cm²)</td>
<td>0.577±0.069</td>
<td>0.561±0.074</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Left hand JSW (mm)</td>
<td>1.68±0.20</td>
<td>1.66±0.21</td>
<td>0.11</td>
</tr>
<tr>
<td>MCP 2 (mm)</td>
<td>1.83±0.23</td>
<td>1.81±0.22</td>
<td>0.15</td>
</tr>
<tr>
<td>MCP 3 (mm)</td>
<td>1.65±0.21</td>
<td>1.63±0.23</td>
<td>0.25</td>
</tr>
<tr>
<td>MCP 4 (mm)</td>
<td>1.55±0.23</td>
<td>1.54±0.24</td>
<td>0.07</td>
</tr>
<tr>
<td>Right hand BMD (g/cm²)</td>
<td>0.589±0.069</td>
<td>0.572±0.075</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Right hand JSW (mm)</td>
<td>1.70±0.19</td>
<td>1.67±0.19</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>MCP 2 (mm)</td>
<td>1.86±0.23</td>
<td>1.83±0.23</td>
<td>0.03</td>
</tr>
<tr>
<td>MCP 3 (mm)</td>
<td>1.66±0.20</td>
<td>1.62±0.21</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>MCP 4 (mm)</td>
<td>1.58±0.23</td>
<td>1.55±0.22</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Both hands JSW (mm)</td>
<td>1.69±0.19</td>
<td>1.66±0.19</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Both hands BMD (g/cm²)</td>
<td>0.583±0.068</td>
<td>0.566±0.074</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Erosion score</td>
<td>1.72±3.26</td>
<td>2.59±3.82</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>JSN Score</td>
<td>2.12±5.08</td>
<td>3.89±6.05</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>SHS</td>
<td>3.84±7.35</td>
<td>6.48±8.22</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

BMD, bone mineral density; JSN, joint space narrowing; JSW, joint space width; MCP, metacarpophalangeal; SHS, Sharp van der Heijde score.

follow-up visits as repeated measures. These models were further adjusted for height, age, gender and BMI, which have been shown to influence JSW.22 25

All tests were bilateral and p values of <0.05 were considered statistically significant. The statistical analyses were performed with SPSS Statistics V.23, Excel 2013 V.15.0 (Microsoft, Redmond, Washington, USA) and Prism V.6.07 for Windows (GraphPad Software, La Jolla, California, USA).

Ethical considerations
All patients gave their informed consent and the study was performed in accordance with the Declaration of Helsinki. Ethical approval for the present study was obtained from the ethical board in all participating units and was registered at http://www.clinicaltrials.gov (NCT00764725).

RESULTS
Baseline patient characteristics
A total of 119 patients with early RA, 78% female and with a mean age of 53 (±14.4) years, were included in this study. Baseline characteristics are shown in table 1, where they are compared with the whole SWEFOT population.

Assessment of JSW
The change of JSW in both hands from baseline to 1 year is shown in table 2 and figure 3A. There is a significant decrease over 12 months in the mean JSW of the two
Assessment of BMD
The detailed baseline and 1 year follow-up BMDs are shown in table 2 and figure 3B. The mean BMD decreased significantly in both hands combined, as well as in the right and left hand separately.

Assessment of modified SHS
SHS and its components, JSN and erosion score, are shown from baseline to 1 year in table 2 and figure 3C. Both the JSN and erosion components increased significantly over 12 months.

Multilevel mixed model
The multilevel model clustering ΔJSW and ΔBMD showed a significant (p<0.0001) positive association from baseline to 1 year follow-up in the 96 patients. This model was adjusted for height, age, gender and BMI.

The JSW for joints with damage (JSN>0) and without damage (JSN=0) was compared using the raw data and a multilevel mixed model (figure 4). The raw data of the mean JSW at baseline and 12 months showed that the undamaged joints were wider than the damaged joints, 1.68 mm (95% CI 1.70 to 1.67) and 1.54 mm (95% CI 1.63 to 1.46), respectively. This value, however, is not statistically significant as it does not account for repeated visits, that is, baseline and follow-up, and that the patient’s own
joints are highly inter-related. The unadjusted multilevel model, however, had accounted for these factors and showed a significant difference in JSW between undamaged and damaged joints, 1.68 mm (95% CI 1.72 to 1.64) vs 1.63 mm (95% CI 1.68 to 1.58) (p=0.0048), and this difference remained significant after adjustment for height, age, gender and BMI, 1.66 mm (95% CI 1.70 to 1.61) vs 1.62 mm (95% CI 1.68 to 1.56) (p=0.042), respectively.

**DISCUSSION**

The aims of this study were to explore and evaluate the changes in JSW and BMD at baseline and 1-year follow-up compared with the JSN component in the SHS. This program is a further development of the program used by Böttcher et al,21 which was a semiautomated program that had an operator-dependent stage of evaluation. The non-dominant hand radiograph was used in the above study, whereas bilateral images of patients with early RA were used in our study. We also incorporated both BMD and JSW using the updated, operator-independent, fully automated program. This operator independence makes it more clinically viable as a diagnostic tool as you introduce less potential operator errors, minimise the time of data acquisition and avoid the training of technicians.

The fully automated method revealed a JSW and BMD reduction between baseline and 1-year follow-up. Significant differences in the JSW between damaged and non-damaged joints as rated by the JSN score were noted. The multilevel mixed model also showed a significant positive association between ∆BMD and ∆JSW from baseline to 1-year follow-up.

It is known that one of the markers of radiographic progression in RA is cartilage damage, displayed as narrowing of the JSW.26−27 Since effective disease-modifying antirheumatic drug (DMARD) treatments may slow down radiographic progression, it is important to be able to detect subtle changes, especially in the early stages of RA, to improve treatment in order to achieve improved radiographic and clinical outcome.28 A sensitive and objective method to detect cartilage loss is valuable, but it also needs to be reproducible. The reproducibility of the fully automated JSW was evaluated with three repeated radiographs of the non-dominant hand of 30 healthy volunteers and resulted in a CV% of 1.4% and an SDD of 0.002 mm. This result is similar to the results presented with the semiautomated program by Pfeil et al.13 High sensitivity and reproducibility are a major strength of automated JSW seen in several studies.7,12,25 In our study, we illustrate that with this automated method, we are able to detect a difference in JSW between damaged and undamaged joints, as defined according to the JSN component of the validated SHS method. Furthermore, the difference in JSW was significant in the unadjusted mixed model and remained significant after adjustment for height, age, gender and BMI, which have previously been shown to be confounding variables for measurements of JSW in RA.22,23,25,29 This suggests that the loss of cartilage as shown by this fully automated JSW measuring method may primarily be due to the progression of the actual disease. Another strength of a fully automated JSW measurement is that it may help remedy the limitations of measuring JSN by SHS or by visual inspection alone. Limitations such as inter-reader and intrareader variations and the necessity of qualified readers are thus eliminated.30 Also important in this study is the need for high-quality acquisition radiographs that are free from technical and positional errors. This for us is a strength rather than a weakness, from the perspective of establishing radiographic quality control and assurance of good radiological practice.

Rapid loss of BMD may take place in the early stages of RA and predicts future radiographic progression as assessed with SHS.10−13 Compared with CR, which reportedly only detects damage if the reduction in bone density is more than 35%,16,17,31 digital BMD measuring has a higher sensitivity. This may be important in the clinic when DMARDs may halt or slow down progression to a point where the CR method is not sensitive enough to measure a difference. Digital BMD measurement, however, is a sensitive tool that has been shown to distinguish treatment effects.13 To measure BMD is an objective and quick method for quantifying bone loss.32 In our study, BMD decreased significantly in both hands during 12 months, suggesting a progression of damage. A significant positive association was observed in the multilevel model between ∆BMD and ∆JSW. This suggests that the loss of BMD and JSW, that is, bone and cartilage loss, are two processes that are occurring concurrently in RA.

In a clinical setting, it is important to have a very low failure rate of image analysis so as to not impede the workflow. In our study the failure rate was 0% for the analysis of six MCP joints per patient — following the
Biomarkers have shown to predict radiographic progressions. Future studies to evaluate the utility of JSW could and bone erosions may also add value to early RA assessment.

Limitations of this study include its design as a post-hoc analysis from the SWEFOT trial, which was not primarily intended for evaluation of JSW or BMD. This resulted in the inclusion of 119 of 487 patients from the original SWEFOT study, of whom only 96 had both baseline and follow-up digital radiographs useful for our evaluation. There were also relatively few patients with progression of joint damage as measured by SHS, since this was a trial with patients with early RA. This however highlighted the utility of a highly sensitive method such as the one used in this study for monitoring early cartilage damage in RA. Another limitation of the study includes the absence of JSW of MCP 1 and 5. The JSW of the proximal interphalangeal (PIP) II–IV joints were available but omitted due to the association of these joints to osteoarthritis (OA). Furthermore, the MCP joints 2 and 3 have a stronger relationship to RA than the PIP joints, which are more prone to be affected by OA. In addition, the PIP joints are also more complicated as they have a bicompartamental configuration, which makes them very sensitive to minor rotations of the hand. Moreover, the unknown characteristics of JSW in RA as the disease progresses through various stages require further investigation. For example, bisphosphonates have shown to prevent generalised bone loss. Their main target is the osteoclast, identified as a potential culprit of focal bone damage caused by inflammatory diseases. The potential effects of bisphosphonates in focal bone damage related to RA are certainly of great interest, and although we have not tested this it may be beneficial in future RA studies to compare BMD and JSW results on patients who are on bisphosphonates compared with those who are not. Although normative values for JSW have been reported previously, further studies on normal cartilage degeneration over time are warranted. Comparative US studies on cartilage damage and bone erosions may also add value to early RA assessments. Future studies to evaluate the utility of JSW could also be done together with the analysis of biomarkers, as biomarkers have shown to predict radiographic progression as measured by SHS.

In summary, to measure the JSW with this fully automated digital program may be useful as a quick and observer-independent tool for evaluating cartilage damage in early RA. We have shown that a fully automated JSW measurement method was associated with the JSW component of the SHS method, and that it had the further clinical feasibility of being quick and observer-independent. We also showed a significant positive association between ΔBMD and ΔJSW from baseline to 1-year follow-up. Fully automated measuring of JSW with the added benefit of BMD makes JSW measuring a potentially useful clinical tool.

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Contributors MP led the writing of the manuscript, analysed data and interpreted results. YK participated in statistical analyses and interpretation of the results. JK was responsible for JSW measuring and sampling data recovery. LA participated in the statistical analyses and interpretation of the results. KF was responsible for the radiographic assessments including the scoring of radiographs. RW inspired the study, designed it and contributed to analysis. All authors were involved in drafting the article or revising it critically for important intellectual content, and all authors approved the final version to be submitted for publication. MP had full access to all of the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

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Competing interests JK is employed by Sectra AB. No other conflicts of interests to declare.

Ethics approval All patients gave their informed consent and the study was performed in accordance with the Declaration of Helsinki. Ethical approval for the present study was obtained from the ethical board in all participating units and was registered (NCT00764725).

Provenance and peer review Not commissioned; externally peer reviewed.

Data sharing statement Access to raw study data can be granted on contact with the main author.

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REFERENCES


