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A feasibility study

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Automatic Quantification of Bone Marrow Edema on MRI of the Wrist in Patients With Early Arthritis: A Feasibility Study

Evgeni Aizenberg,1* Edgar A.H. Roex,1,2 Wouter P. Nieuwenhuis,3 Lukas Mangnus,3 Annette H.M. van der Helm-van Mil,3 Monique Reijnierse,1 Johan L. Bloem,1 Boudewijn P.F. Lelieveldt,1,4 and Berend C. Stoel1

Purpose: To investigate the feasibility of automatic quantification of bone marrow edema (BME) on MRI of the wrist in patients with early arthritis.

Methods: For 485 early arthritis patients (clinically confirmed arthritis of one or more joints, symptoms for less than 2 years), MR scans of the wrist were processed in three automatic stages. First, super-resolution reconstruction was applied to fuse coronal and axial scans into a single high-resolution 3D image. Next, the carpal bones were located and delineated using atlas-based segmentation. Finally, the extent of BME within each bone was quantified by identifying image intensity values characteristic of BME by fuzzy clustering and measuring the fraction of voxels with these characteristic intensities within each bone. Correlation with visual BME scores was assessed through Pearson correlation coefficient.

Results: Pearson correlation between quantitative and visual BME scores across 485 patients was $r=0.83$, $P<0.001$.

Conclusions: Quantitative measurement of BME on MRI of the wrist has the potential to provide a feasible alternative to visual scoring. Complete automation requires automatic detection and compensation of acquisition artifacts. Mag Reson Med 79:1127–1134, 2018. © 2017 The Authors Magnetic Resonance in Medicine published by Wiley Periodicals, Inc. on behalf of International Society for Magnetic Resonance in Medicine. This is an open access article under the terms of the Creative Commons Attribution-NonCommercial-NoDerivs License, which permits use and distribution in any medium, provided the original work is properly cited, the use is non-commercial and no modifications or adaptations are made.

INTRODUCTION

The presence of bone marrow edema–like abnormalities (BME) has been shown to be a strong predictor of radiographic progression in rheumatoid arthritis (RA) patients (1–4), and is therefore an important biomarker in early arthritis. Evaluation of BME is done on MRI, where it is visually scored based on the Outcome Measures in Rheumatology RA-MRI Scoring (RAMRIS) system (5,6). This scoring method requires a trained reader to visually estimate the volume of BME. Such estimates are challenging and time-consuming because of the need to assess multiple imaging planes, and slices and are inherently undermined by the simultaneous contrast effect (7–9) of the human visual system, which causes the reader to perceive the same image intensity value differently, depending on surrounding background intensities.

An automatic and quantitative approach to evaluating BME on MR scans could overcome the limitations of visual scoring by offering high-precision measurements derived directly from 3D image data. It could alleviate the time burden of training and manual scoring for clinical researchers, and could facilitate the use of MRI in drug evaluation studies, in which using a trained team of readers is costly.

Several previous studies on BME quantification in the wrist joint (10–12) relied on a semi-automatic method proposed by Li et al. (13). However, this technique requires an expert to manually delineate non-edema and edema regions of interest within every bone that needs to be evaluated. These studies were also limited to a small sample size of fewer than 20 subjects. One related study focuses on fully automatic BME quantification in the knee joint (14), but it is not directly clear how to extend the bone segmentation method (15) to a joint with more than two bones, as is the case in the wrist.

In the work presented here, we developed an automatic framework for measuring the fraction of bone volume affected by BME in the eight carpal bones of the wrist joint. In contrast to previous methods, we used atlas-based segmentation to automatically locate and...
delineate the carpal bones. Our aim was to investigate the feasibility of BME quantification through such atlas-based approach, and assess the correlation between quantitative measurements and visual BME scores in a large cohort of early arthritis patients.

**METHODS**

**Patients**

A total of 573 early arthritis patients from the Leiden Early Arthritis Clinic cohort (16) (mean age, 54.7 years; age range, 18.1–87.9 years) were studied: 354 female (mean age, 53.0 years; age range, 18.7–85.3 years) and 219 male (mean age, 57.5 years; age range, 18.1–87.9 years) patients. Inclusion required clinically confirmed arthritis by physical examination in one or more joints, and symptom duration of less than 2 years. The MR scans were obtained for the wrist joint of the most painful side (or the dominant side in cases of equally severe symptoms on both sides). The study was approved by the institutional medical ethics committee and all participants provided written informed consent.

**MRI Sequences**

The wrist joint was scanned with an ONI MSK Extreme 1.5 Tesla (T) extremity MR scanner (GE Healthcare, Waukesha, WI) with a 100-mm coil. Before contrast agent injection, T1-weighted fast spin-echo sequence (T1) was acquired in the coronal plane with repetition time (TR) of 650 ms, echo time (TE) of 17 ms, acquisition matrix 388 × 88, echo train length (ETL) 2. After intravenous injection of Gd-chelate (gadoteric acid, Guerbet, Paris, France, standard dose of 0.1 mmol/kg), a T1-weighted fast spin-echo sequence with frequency-selective fat saturation (T1-Gd) was obtained in the coronal plane (TR 650 ms/TE 17 ms, acquisition matrix 364 × 224, ETL 2) and the axial plane (TR 570 ms/TE 7 ms, acquisition matrix 320 × 192, ETL 2). Coronal sequences were acquired with a slice thickness of 2 mm and a slice gap of 0.2 mm. Axial sequences were acquired with a slice thickness of 3 mm and a slice gap of 0.3 mm. The use of a T1-Gd sequence instead of a T2-weighted fat-saturated sequence is a validated modification that has been shown to permit equally well in the depiction of BME and allows for a faster scanning protocol (17,18), which in turn reduces patient discomfort. Safety risk was minimized to the degree possible by the use of a macrocyclic contrast agent (19,20).

**Visual Scoring of BME**

Bone marrow edema was assessed in line with the definitions proposed by RAMRIS (5) with validated modification of substituting T2-weighted fat-saturated sequence with T1-Gd sequence (17,18). The BME was independently scored by two trained readers who were blinded to clinical data on a 0–3 scale based on the estimated fraction of affected bone volume: 0, no BME; 1, 1–33% of bone edematous; 2, 34–66%; 3, 67–100%. The within-reader intraclass correlation coefficients (ICCs) for the total inflammation score were 0.98 and 0.93; the between-reader ICC was 0.95. The mean BME score of the two readers was considered.

Patients for which at least one reader marked one or more bones as unscorable (typically a result of fat suppression issues) were excluded (n=11). Patients whose T1-Gd images suffered from incomplete fat suppression, but still considered scorable by readers based on T1 images showing low signal intensity in the matching areas with BME on T1-Gd, were retained.

**Quantitative Image Analysis Framework**

Our automatic framework consisted of three stages. First, super-resolution reconstruction was applied to fuse coronal and axial T1-Gd scans into a single high-resolution 3D image. Next, the carpal bones were located and delineated using atlas-based segmentation. Finally, the extent of BME within each bone was quantified by identifying image intensity values characteristic of BME by fuzzy clustering, and measuring the fraction of voxels with these characteristic intensities within each bone. Note that because the super-resolution reconstruction step requires a coronal and axial scan of the same sequence as input, this stage, and therefore the entire framework, could only be applied to T1-Gd scans. Therefore, pre-contrast T1 images, which were acquired only in the coronal plane, were not used in the quantitative image analysis framework.

**Super-resolution Reconstruction**

When readers evaluate BME visually, they make use of two complementary scans: one acquired in the coronal plane and the second in the axial plane. This is because the slice thickness in each of the scans (2 mm in coronal; 3 mm in axial) is much larger than the in-plane spacing between voxels (~0.2 mm). Therefore, one scan compensates for anatomical detail lost in the other scan, allowing the reader to perceptually form a more complete assessment of the anatomy. Naturally, this raises the question of how to simulate such perceptual fusion of two images on the computer, to obtain a single 3D image with isotropic voxels and high resolution in all three viewing planes. This type of problem (i.e., reconstruction of a high-resolution image of an object from multiple low-resolution images of the same object) is commonly referred to as super-resolution reconstruction (SRR).

A variety of SRR methods have been proposed for MRI (21–24). In this study, we applied the method developed by Poot et al. (24). This algorithm belongs to the family of spatial domain SRR methods, which construct a linear model of the image acquisition system and reconstruct the high-resolution image by solving a system of linear equations. This system is often underdetermined, as in our case, and is solved by applying regularization. We used Laplacian regularization with parameter λ = 0.05. This value was optimized in an experiment by two expert radiologists (M.R. and J.L.B.) to provide satisfactory balance between image noise/artifacts and visual clarity of BME, synovial tissue, cartilage, and fluid around tendons. Before applying SRR, the axial scan was spatially aligned to the coronal scan using the Elastix
software package (25,26), axial image intensity was linearly matched to the coronal image intensity, and the field of view of both images was restricted to the overlapping physical space between the two scans. Figure 1 shows an example of applying SRR to a pair of coronal and axial scans.

### Segmentation of Carpal Bones

The carpal bones were located and delineated using atlas-based segmentation (ABS) (27). The atlas consisted of 13 early arthritis patients. For each atlas patient, the carpal bones were manually segmented in the coronal and axial T1-Gd images, yielding two segmentation images. The voxels of these manual segmentation images were assigned an integer bone label value ranging from 1 to 8 in locations corresponding to one of the eight carpal bones, or otherwise the value 0 in locations outside the bones. Then, separately for each bone, the two manual segmentation images were fused using SRR. Voxels with values above 78% of the bone label value were assigned the bone label value, and the remaining voxels were zeroed to discard noise. The resulting eight images were superimposed to obtain the complete segmentation image in high-resolution space.

The first phase of the ABS routine consisted of image registration between each of the 13 atlas images and the target image being segmented. Image registration (using Elastix (25)) was done in two stages (28): first, a similarity mapping to account for global translation, rotation and scaling, followed by a B-spline mapping to account for local deformations. After spatially mapping carpal bone segmentations from every atlas image onto the target image, a majority vote was applied across all mappings, determining whether a voxel was labeled as background or as one of the carpal bones.

It should be noted that all atlas images contained the right wrist joint. For segmentation of the left wrist, atlas images were horizontally mirrored before registration. To avoid biased measurements, patients that were part of the ABS atlas were excluded from optimization and validation phases.

<table>
<thead>
<tr>
<th>Coronal plane</th>
<th>Axial plane</th>
<th>Sagittal plane</th>
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</thead>
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<tr>
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<td><img src="image2" alt="Axial image" /></td>
<td><img src="image3" alt="Sagittal image" /></td>
</tr>
<tr>
<td><img src="image4" alt="Super-resolution image" /></td>
<td><img src="image5" alt="Super-resolution image" /></td>
<td><img src="image6" alt="Super-resolution image" /></td>
</tr>
</tbody>
</table>

**FIG. 1.** Coronal, axial, and super-resolution images (top to bottom rows, respectively) and their coronal, axial, and sagittal viewing planes (left to right columns, respectively). The original scans exhibit high resolution only in one plane, whereas the super-resolution image exhibits high resolution in all three planes.
Assessment of ABS Accuracy

To assess the accuracy of ABS, a leave-one-out cross-validation was performed. In each of the 13 runs, 12 out of 13 atlas images would constitute the atlas set, and the remaining image would be used as the target image to be segmented. The result was validated against manual segmentation of the coronal T1-Gd image. Segmentation accuracy was evaluated by computing precision and recall rates for each carpal bone. Here, “precision rate” refers to the fraction of voxels segmented by ABS that overlap with the manual bone segmentation, whereas “recall rate” refers to the fraction of voxels within the manual bone segmentation that were correctly segmented by ABS.

Bone Marrow Edema Quantification

Bone marrow edema is characterized by high signal intensity on T1-Gd images as a result of contrast enhancement and the suppressed normal fatty bone marrow. The precise intensity values vary per acquisition, depending on the strength of contrast enhancement and fat suppression. The variation of these values is further broadened by inherent magnetic field inhomogeneities of the MR scanner. To account for these acquisition-specific intensity ranges of edematous versus non-edematous bone marrow, fuzzy C-means clustering (29,30) was applied to the intensity values of all voxels in each image, assuming two clusters. This yields two probability map images (one per cluster), in which each voxel contains the probability of that voxel belonging to the respective cluster. Let $C_2$ be the cluster whose center value is the higher of the two computed cluster centers.

As Figure 2 illustrates, high probabilities (bright voxels) within the $C_2$ probability map correspond to locations of high fluid content, such as BME and synovium.

For each carpal bone, the fraction of bone affected by BME was estimated as the fraction of voxels (out of the total number of voxels within the bone’s segmentation) whose probability of belonging to $C_2$ was higher than the threshold value $T_{C_2}$ (numeric value optimized below). The resulting quantitative BME measurement (BME-QM) takes any fractional values between 0 and 1.

Optimization

To optimize the $T_{C_2}$ threshold parameter based on correlation with visual BME scores, a training set of patients was defined. The number of patients with low-moderate BME in our cohort is much larger than the number of patients with severe BME. Therefore, random sampling

<table>
<thead>
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<th>Patient category index</th>
<th>$R_{max}$ interval</th>
<th>Number of patients</th>
</tr>
</thead>
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<td>0</td>
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<td>189</td>
</tr>
<tr>
<td>1</td>
<td>$0 &lt; R_{max} \leq 1$</td>
<td>208</td>
</tr>
<tr>
<td>2</td>
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<td>42</td>
</tr>
<tr>
<td>3</td>
<td>$2 &lt; R_{max} \leq 3$</td>
<td>29</td>
</tr>
</tbody>
</table>

Note: Random sampling across all categories would form a training set that consists primarily of patients with $R_{max} \leq 1$. In contrast, randomly selecting 15 patients from category 3, for example, guarantees that the training set will include 15 patients, with at least one bone that received a visual BME score greater than 2. Thus, random sampling from individual categories helps to ensure that $T_{C_2}$ is optimized with respect to the entire range of the visual BME score.
of the cohort does not guarantee inclusion of patients with severe BME in the population sample. To ensure that patients with a high degree of BME were represented in the training set, we categorized a set of 468 patients by the maximum visual BME score ($R_{\text{max}}$) across the carpal bones. Four sampling categories were defined corresponding to four intervals within $R_{\text{max}}$ range. Table 1 lists the defined categories and the number of patients that fall into each category. Next, 15 patients were randomly selected from each category to form a training set of 60 patients.

To avoid training errors, three patients whose MR scans suffered from incomplete fat suppression and one patient for which ABS failed were excluded from the obtained training set. This brought the final training set size to 56 patients. The optimal value of $TC_2$ was found by maximizing the Pearson correlation coefficient $r$ between the sum of visual BME scores across all carpal bones and the sum of BME-QM across all carpal bones.

**Validation**

After optimizing and locking the value of $TC_2$, the method was validated on 502 patients who were not part of the training set.

**Statistical Analysis**

The Pearson correlation coefficient $r$ between the sum of visual BME scores across all carpal bones and the sum of BME-QM across all carpal bones was evaluated. $P$ values below 0.05 were indicative of statistical significance. The MR scans that suffered from incomplete fat suppression were noted and excluded from the correlation computation. Scans with other acquisition artifacts, such as noise patterns and incomplete field of view, were excluded from the analysis. Patients in which one or more bones were not segmented by ABS yielded undefined values for BME-QM. Because undefined values cannot be included in the correlation computation, these patients were excluded from statistical analysis. The statistics were computed using MATLAB R2015b (The MathWorks Inc, Natick, MA).

**RESULTS**

**Assessment of ABS Accuracy**

The mean bone-level recall and precision rates of ABS with respect to manual segmentations across 13 patients are shown in Figure 3. Recall rates were lowest in the pisiform (mean of 0.58 ± 0.09 SD) and highest in the capitate (mean of 0.82 ± 0.03 SD). Precision rates were high in all bones, with mean values ranging from 0.92 to 0.96 and SD values ranging from 0.02 to 0.05.

**Optimization**

The maximum Pearson correlation ($r = 0.86, P < 0.001$), over 56 training set patients, between the sum of visual BME scores across all carpal bones and the sum of BME-QM across all carpal bones, was achieved at threshold value $T_{C2} = 0.83$ (Figure 4). The scatter plot of the data is shown in Figure 5.

**Validation**

Out of 502 patients, BME-QM was undefined in six patients because of failed segmentation. Three patients...
were excluded as a result of noise artifacts \((n=2)\) and incomplete field of view \((n=1)\) in their images. The MR scans of eight patients suffered from incomplete fat suppression. For the remaining 485 patients, the Pearson correlation between the sum of visual BME scores across all carpal bones and the sum of BME-QM across all carpal bones was \(r = 0.83, P < 0.001\). The scatter plot of the data is shown in Figure 6. Most patients formed clusters of steadily increasing BME-QM values, as the visual score value increased. Some outliers from this general trend were clearly visible for visual score value of 0 and BME-QM values between 1 and 2. These high quantitative values were the result of inaccurate segmentation of the carpal bones. Several patients whose images suffered from incomplete fat suppression produced BME-QM values that were largely deviating from the observed regression fit.

**DISCUSSION AND CONCLUSIONS**

In this study, we investigated the feasibility of automatic quantification of BME on MRI of the wrist in patients with early arthritis through an atlas-based approach. We chose to focus on the carpal bones, as they provide a complex multi-object scenario for exploring the feasibility of an atlas-based quantification framework. The advantage of this framework is that it can be straightforwardly expanded to other areas of the wrist and other joints by adding these areas of interest to the atlas. Validation results across 485 early arthritis patients indicated good correlation between BME-QM and visual BME scores. It should be noted that perfect correlation is inherently not achievable because of the coarse grading scale of the visual score and the fine grading scale of BME-QM.

Our training strategy helped to ensure that during validation BME-QM correlated well across the entire range of the visual BME score. The fact that the correlation curve in Figure 4 is relatively flat for \(T_{C2}\) values between 0.75 and 0.9 suggests that there is a range of \(T_{C2}\) values in this interval that result in good agreement between quantitative and visual scores. Furthermore, because BME-QM measures the fraction of voxels with \(C2\) probability above \(T_{C2}\), this appears to indicate that locations considered as BME in visual scoring often result in \(C2\) probability values around 0.9. We also examined the effect of a smaller training set on \(T_{C2}\) optimization (data not shown), with five patients randomly selected from each \(R_{max}\) category forming a training set of 20 patients. We observed a similarly stable high correlation for \(T_{C2}\) values between 0.75 and 0.9, suggesting that the optimization step is not overly sensitive to training set size, as long as patients from all categories of BME severity \((R_{max})\) are represented in the training data.

The time required to execute the BME-QM framework for one patient on an Intel Xeon E5-1620 v3 CPU was approximately 58 min (SRR, ~20 min; ABS, ~35 min; BME quantification ~3 min). Atlas-based segmentation is the most time-consuming step, but it can be accelerated 10-fold by running image registrations between all atlas images and the target image in parallel. Because registrations are independent of each other, this can be easily achieved given sufficient computing power. However, in large cohort studies, in which evaluation of image data is often carried out days or weeks after the image is acquired, such acceleration may be irrelevant; an automatic framework can be executed immediately after image acquisition in an integrated fashion, thus ensuring quantitative results are available by the time a research project enters the evaluation phase.

Atlas-based segmentation provided satisfactory segmentation for the vast majority of patients. In practice, failed segmentation cases will require manual

![FIG. 5. Scatter plot of sum of BME-QM across all carpal bones versus sum of visual BME scores across all carpal bones for 56 training set patients. Each data point represents a single patient. \(r = 0.86, P < 0.001, T_{C2} = 0.83\). Dashed black line represents linear regression fit.](image)

![FIG. 6. Scatter plot of sum of BME-QM across all carpal bones versus sum of visual BME scores across all carpal bones for 493 validation set patients. Each data point represents a single patient. Linear regression fit (dashed black line) and Pearson correlation \(r\) were computed over 485 patients whose MR scans did not suffer from incomplete fat suppression (circular data points): \(r = 0.83, P < 0.001, T_{C2} = 0.83\).](image)
adjustment by an expert in order for BME-QM to be computed. Oversegmentation of bones or shifted segmentations that include synovium voxels increase the value of BME-QM as a result of contrast enhancement in the synovium. It is preferable to slightly undersegment the bone to ensure the exclusion of synovium while retaining most of the bone marrow within the segmentation. That said, significant undersegmentation may lead to an upward bias in BME-QM. Quantitative assessment of ABS accuracy in 13 patients revealed the tendency of ABS to undersegment bones (midrange recall rates and high precision rates). Therefore, the current framework may raise false alarms when bone volume is underestimated in the presence of moderate BME. The midrange recall rates also suggest unwanted variability in BME-QM as a result of incomplete bone segmentation. The fact that the lowest recall rates were observed in the pisiform while the highest in the capitate, is likely because the pisiform is the smallest of the carpal bones, whereas the capitate is the largest of the carpal bones. An additional challenge during registration is the varied intensity and pattern of BME across patients. It is therefore advisable to avoid using very fine grid spacing during the B-spline registration step, as alignment between images on a coarser scale should be less sensitive to these local variations. Another potential pitfall is inclusion of erosions in the segmentation result. Erosions may contain high intensities that will mistakenly contribute to the value of BME-QM. To address these possible pitfalls and improve bone-level recall rates, an automatic refinement step should follow ABS in the future. In addition, to ensure robustness of the atlas to variations in MRI acquisition protocols and scanners at different sites, it may be necessary to form a larger atlas set consisting of subatlases of wrist scans acquired under different echo/repetition times and magnetic field strengths. The most suitable subatlas can then be automatically identified based on the acquisition parameters of a specific target image.

Incomplete fat suppression during acquisition of MR scans has an adverse effect on the accuracy of BME-QM. Bone marrow fat signal that is not properly suppressed results in high intensities that are mistaken for edema voxels by the clustering algorithm. Fat suppression quality requirements for BME-QM are higher compared with visual scoring. This is because of the availability of pre-contrast image data in visual scoring and pattern recognition during visual assessment of increased signal intensity secondary to insufficient fat suppression. Although fat suppression issues are relatively rare, they must be identified before applying BME-QM to reduce false positives. The possibility of identifying and compensating for fat suppression issues automatically should be investigated. In addition, more robust fat suppression techniques that are less sensitive to bulk susceptibility, such as Dixon techniques, may be beneficial when BME-QM is used.

A limitation of the current study is that the quality of carpal bone segmentation in training and validation set patients was judged subjectively. Quantitative assessment of segmentation accuracy was not possible, as no ground truth, manual segmentations were available for these patients. Quantification of segmentation accuracy would allow to supplement the BME measurement with a confidence measure. Another limitation is that pre-contrast image data could not be included in the framework, as pre-contrast T1 scans were acquired only in the coronal plane, whereas SRR requires a coronal and axial scan of the same sequence as input. Therefore, a straightforward voxel-to-voxel comparison between SRR T1-Gd images and pre-contrast T1 images was not possible. Inclusion of pre-contrast data would allow to explore a subtraction methodology as a means of quantifying BME, and could facilitate the detection of fat suppression issues.

Recently, another framework aimed at automatically quantifying RA-related biomarkers, called quantitative RAMRIS, was proposed by Bowes et al. (31,32) and used in a treatment effects study by Conaghan et al. (33). These studies focus on measuring change over time, demonstrating higher sensitivity of quantitative measurements compared with RAMRIS. In contrast, we focused on validation of quantitative measurements at a single time point. In the future, it would be interesting to use BME-QM for measuring change over time and evaluate its sensitivity.

We conclude that BME-QM has the potential to provide a feasible alternative to visual scoring of BME on MRI of the wrist in patients with early arthritis. Complete automation requires further refinement of carpal bone segmentation and automatic detection and compensation of acquisition artifacts. Future work should also add more locations of interest relevant to RA to the atlas and extend this framework to other types of inflammation, such as synovitis and tenosynovitis. These developments can save time and manual effort for clinical researchers and help assess the value of MRI both for diagnosing RA and monitoring its treatment.

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