Assessment of inflammatory activity in rheumatoid arthritis: a comparative study of clinical evaluation with grey scale and power Doppler ultrasonography

E Naredo, G Bonilla, F Gamero, J Uson, L Carmona, A Laffon


Objective: To compare the clinical assessment of overall inflammatory activity in patients with rheumatoid arthritis (RA) with grey scale and power Doppler (PD) ultrasonography (US).

Methods: Ninety four consecutive patients with RA were included. Demographic and clinical data, C reactive protein (CRP) level, and erythrocyte sedimentation rate (ESR) were recorded for each patient. The presence of tenderness, swelling, and a subjective swelling score from 1 to 3 were independently assessed by two rheumatologists, who reached a consensus in 60 joints examined in each patient. All patients underwent a US examination by a third blinded rheumatologist, using PD. US joint effusion, synovitis, and PD signal were graded from 1 to 3 in the 60 joints. Joint count and joint index for effusion, synovitis, and PD signal were recorded. A 28 joint count for clinical and US variables was calculated. Interobserver reliability of the US examination was evaluated by a fourth blinded rheumatologist.

Results: US showed significantly more joints with effusion (mean 15.2) and synovitis (mean 14.6) than clinical examination (mean 11.5, p<0.05). A significant correlation was found between joint count and joint index for swelling, US effusion, synovitis, and PD signal. The 28 joint count for effusion, synovitis, and PD signal correlated highly with the corresponding 60 joint counts. US findings correlated better with CRP and ESR than clinical measures. Interobserver reliability was better for US findings than for clinical assessment.

Conclusion: US is a sensitive method for assessing joint inflammatory activity in RA, complementary to clinical evaluation.

Rheumatoid arthritis (RA) is a chronic inflammatory disease characterised by the development of synovitis, which damages cartilage, bone, ligaments, and tendons. Assessment of inflammatory activity is essential in daily practice to enable therapeutic decisions and to evaluate disease outcome and response to treatment.

Traditionally, the degree of disease activity has been evaluated by measuring subjective clinical variables, laboratory measures, and radiographic findings. However, clinical evaluation of joint pain and swelling have not been sufficiently reliable, and conventional plain radiography depicts indirect signs of cartilage loss and bony erosions due to previous destructive synovial inflammatory activity.

High frequency ultrasonography (US) has greatly improved musculoskeletal imaging in rheumatology. Several studies have demonstrated that high frequency US is accurate for detecting joint effusion and synovitis, compared with magnetic resonance imaging (MRI) and direct arthroscopic visualisation. US is more sensitive and reproducible than clinical evaluation in assessing joint inflammatory activity.

Power Doppler (PD) US is a new technique of colour Doppler that improves the sensitivity to detect flow from small vessels and low velocity flow at the microvascular level. PD US detects indirect signs of increased vascularisation associated with soft tissue musculoskeletal inflammatory and infectious diseases and enthesitis in spondyloarthopathies. The PD signal correlates highly with local clinical evaluation of joint inflammatory activity in the knee, metacarpophalangeal (MCP) and interphalangeal joints of patients with RA and other inflammatory arthropathies. Recent studies have shown that PD synovial vascularity correlates highly with histologically proved knee pannus and with the degree of synovial vascularisation of the knee and hip.

This study aimed at comparing grey scale US and PD US with clinical and biological findings in the determination of global inflammatory activity assessed in 60 joints of patients with RA. To the best of our knowledge, this is the first US and clinical study that has examined so many joints in each patient with RA.

PATIENTS AND METHODS

Ninety four consecutive patients (20 male, 74 female) who fulfilled the 1987 American Rheumatism Association criteria for RA attending the outpatient rheumatology clinic were included. Mean (SD) age was 57.6 (14.3) years (range 23–88) and mean (SD) disease duration was 69.3 (58.2) months (range 5–280). Patients who had had traumatic, septic, or microcrystalline arthritis, previous joint surgery, or isotopic synovectomy within the past 12 months before the study were excluded.

The following data were recorded for each patient: age, sex, disease duration, drugs received for RA at entry, rheumatoid factor (measured by nephelometry, normal level 0–20 IU/ml), and previous joint surgery or isotopic synovectomy. C reactive protein (CRP) level (measured by nephelometry, normal range 0–5 mg/l) and erythrocyte sedimentation rate (ESR).

Abbreviations: CRP, C reactive protein; ESR, erythrocyte sedimentation rate; HAQ, Health Assessment Questionnaire; MCP, metacarpophalangeal; MRI, magnetic resonance imaging; PD, power Doppler; PIP, proximal interphalangeal; PRF, pulse repetition frequency; RA, rheumatoid arthritis; SJC, swollen joint count; SJ, swollen joint index; TJC, tender joint count; US, ultrasonography; VASQA, visual analogue scale for patient overall assessment of disease activity; VASP, global pain intensity visual analogue scale.
measured by the Westergren method, VESMATIC 60, version 2.05; Menarini Laboratory, Barcelona, Spain) were recorded from each patient’s routine laboratory test performed within 1 week of the study. Informed consent was obtained from all patients before the clinical and US evaluation.

Clinical assessment
The clinical evaluation was performed independently and sequentially by two blinded rheumatologists. One week before the study they carried out a consensus joint examination in patients with RA (not included in the study) for 20 hours. The following bilateral joints were assessed for tenderness and swelling: glenohumeral, acromioclavicular, sternoclavicular, elbow, wrist, MCP, proximal interphalangeal (PIP) of hands, knee, ankle, subtalar, mid-tarsal, metatarsophalangeal, and PIP of feet (total in 94 patients 5640 joints). Hip joints were assessed for tenderness and pain on passive motion. Hip swelling was indirectly considered if pain on passive motion was detected by physical examination. A subjective score from 1 to 3 was assigned for all swollen joints except for the hip (1 = mild; 2 moderate; 3 = marked). Immediately after physical examination, GB and FG compared their findings. If there were discrepancies for the presence or absence of joint tenderness and swelling or the swollen joint scores, they carried out a third examination together to reach consensus. These last results were compared with US findings. Individual physical examinations were used for estimating clinical interobserver agreement for tenderness, swelling and swelling scores. Tender joint count (TJC), swollen joint count (SJC), and a 60 swollen joint index (SJI; sum of the swelling score from each joint) were recorded for each patient.

US examination
All patients underwent a US examination within 30 minutes of the clinical evaluation by a single rheumatologist experienced in US (EN) who was unaware of the clinical findings. US examination was performed with two commercially available ultrasound real time scanners (Logiq 400CL, Table 1

<table>
<thead>
<tr>
<th>Joint</th>
<th>Criteria of effusion/synovitis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glenohumeral</td>
<td>Posterior recess, transducer transverse to the humerus, shoulder in neutral position: maximum distance from the posterior labrum to the posterior infraspinatus and teres minor tendon (posterior capsule) &gt;3 mm</td>
</tr>
<tr>
<td>Acromioclavicular and sternoclavicular</td>
<td>Transducer longitudinal to the joint: maximum distance from the articular bony margin to the joint capsule &gt;3 mm</td>
</tr>
<tr>
<td>Elbow</td>
<td>Longitudinally from the anterior recess with the joint in extension: maximum distance from the humeral capitulum or the coronoid fossa to the joint capsule &gt;2 mm</td>
</tr>
<tr>
<td>Wrist</td>
<td>Longitudinally from the dorsal aspect with the joint in neutral position: maximum distance from the bones to the joint capsule &gt;2 mm</td>
</tr>
<tr>
<td>MCP and PIP joints of hands</td>
<td>Longitudinally from the dorsal view with the joint in extension: maximum distance from the articular bony margin to the joint capsule &gt;2 mm</td>
</tr>
<tr>
<td>Hip</td>
<td>Anterior recess, transducer longitudinal to the femoral neck, with the joint in neutral position: maximum distance from the cortex, at the middle of the concavity of the femoral neck to the joint capsule &gt;7 mm or loss of concavity of the joint capsule</td>
</tr>
<tr>
<td>Knee</td>
<td>Longitudinally from the suprapatellar recess, in a supine position, with the joint in 30˚ of flexion: maximum anterior-posterior diameter of the suprapatellar bursa &gt;3 mm</td>
</tr>
<tr>
<td>Ankle</td>
<td>Longitudinally from the anterior recess with the joint in slight plantar flexion: maximum distance from the talus bone to the joint capsule &gt;2 mm</td>
</tr>
<tr>
<td>Subtalar</td>
<td>Longitudinally to the joint with the ankle in slight plantar flexion: maximum distance from the articular bony margin to the joint capsule &gt;2 mm</td>
</tr>
<tr>
<td>Mid-tarsal</td>
<td>Longitudinally from the dorsal view: maximum distance from the articular bony margin to the joint capsule &gt;2 mm</td>
</tr>
<tr>
<td>MTP and PIP of feet</td>
<td>Longitudinally from the dorsal view with the joints in extension: maximum distance from the articular bony margin to the joint capsule &gt;2 mm</td>
</tr>
</tbody>
</table>

Figure 1 Longitudinal sonographic image of the wrist joint with moderate effusion, moderate synovitis, and mild (A), moderate (B), and marked (C) colour signal.
subjectively graded from 1 to 3 (1 = mild; 2 = moderate; 3 = marked).

Joint effusion and synovitis were defined as echogenic non-compressible intra-articular tissue, articular material, within synovial recesses. Synovitis was defined as hypoechoic or anechoic compressible intra-articular material, within synovial recesses. Synovitis was subjectively graded from 1 to 3 (1 = mild; 2 = moderate; 3 = marked).

Synovial vascularisation was assessed by PD US in each of the 60 joints. PD imaging was performed by selecting a region of interest that included the bony margins, articular space, and a variable view of surrounding tissues (depending on the joint size). PD variables were adjusted to the lowest permissible pulse repetition frequency (PRF) to maximise sensitivity. This setting resulted in a PRF between 500 and 1000 Hz depending on the joint scanned. Low wall filters were used. The dynamic range was 20–40 dB. Colour gain was set just below the level at which colour noise appeared underlying bone (no flow should be visualised at the bony surface). This setting resulted in gains of from 18 to 30 dB. An image with maximal colour activity was selected for analysis. Flow was additionally demonstrated in two planes and confirmed by pulsed wave Doppler spectrum to exclude artefacts.

The intra-articular PD signal was subjectively graded on a semiquantitative scale from 0 to 3 (0 = absence, no intra-articular flow; 1 = mild, single vessel signal; 2 = moderate, confluent vessels; 3 = marked, vessel signals in more than half of the intra-articular area). In each patient we recorded joint count for US effusion (USJCE), joint count for synovitis (USJCS), joint count for PD signal (USJCPD), and a 60 joint index for effusion (USJIE), synovitis (USJSI), and PD signal (USJIPD) (sum of the effusion, synovitis, and PD signal scores, respectively, obtained from each joint). In addition, we calculated for each patient a 28 joint count (10 PIP joints and 18 DIP joints) (USJIPD) (sum of the effusion, synovitis, and PD signal scores, respectively, obtained from each joint).

Table 2  Demographic, clinical, and laboratory characteristics of patients evaluated with scanners 1 and 2

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Scanner 1</th>
<th>Scanner 2</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age (years)</td>
<td>58.1 (14.3)</td>
<td>56.3 (14.3)</td>
<td>NS</td>
</tr>
<tr>
<td>Disease duration (months)</td>
<td>68.3 (57.9)</td>
<td>72.2 (60.1)</td>
<td>NS</td>
</tr>
<tr>
<td>Pain by 0–100 VAS (mm)</td>
<td>30.5 (26)</td>
<td>36.3 (30.3)</td>
<td>NS</td>
</tr>
<tr>
<td>Patient’s overall assessment by 0–100 VAS (mm)</td>
<td>38.2 (26.1)</td>
<td>38.3 (28.6)</td>
<td>NS</td>
</tr>
<tr>
<td>Tender joint count</td>
<td>14 (16.1)</td>
<td>9.8 (13)</td>
<td>NS</td>
</tr>
<tr>
<td>Swollen joint count</td>
<td>10.9 (7.8)</td>
<td>13.2 (6.3)</td>
<td>NS</td>
</tr>
<tr>
<td>Swollen joint index</td>
<td>69.8 (9.7)</td>
<td>72.6 (8.4)</td>
<td>NS</td>
</tr>
<tr>
<td>Health Assessment Questionnaire score (0–3)</td>
<td>1 (0.7)</td>
<td>0.9 (0.7)</td>
<td>NS</td>
</tr>
<tr>
<td>Erythrocyte sedimentation rate (mm/1st h)</td>
<td>28.4 (19.4)</td>
<td>25.9 (14.5)</td>
<td>NS</td>
</tr>
<tr>
<td>C reactive protein (mg/l)</td>
<td>15 (19)</td>
<td>16 (15)</td>
<td>NS</td>
</tr>
</tbody>
</table>

Results are expressed as mean (standard deviation).

VAS, visual analogue scale; NS, non-significant.

Table 3  Correlation between clinical, ultrasonographic, and laboratory variables

<table>
<thead>
<tr>
<th>HAQ</th>
<th>VASP</th>
<th>VASOA</th>
<th>ESR</th>
<th>CRP</th>
</tr>
</thead>
<tbody>
<tr>
<td>TJC</td>
<td>0.40*</td>
<td>0.46*</td>
<td>0.36*</td>
<td>0.12</td>
</tr>
<tr>
<td>SJC</td>
<td>0.44*</td>
<td>0.41*</td>
<td>0.33*</td>
<td>0.41*</td>
</tr>
<tr>
<td>SJI</td>
<td>0.42*</td>
<td>0.39*</td>
<td>0.32*</td>
<td>0.45*</td>
</tr>
<tr>
<td>TJC28</td>
<td>0.42*</td>
<td>0.48*</td>
<td>0.39*</td>
<td>0.10</td>
</tr>
<tr>
<td>SJC28</td>
<td>0.38*</td>
<td>0.35*</td>
<td>0.29*</td>
<td>0.39*</td>
</tr>
<tr>
<td>USJCE</td>
<td>0.13</td>
<td>0.23*</td>
<td>0.31*</td>
<td>0.50*</td>
</tr>
<tr>
<td>USJCS</td>
<td>0.15</td>
<td>0.26*</td>
<td>0.32*</td>
<td>0.50*</td>
</tr>
<tr>
<td>USJCPD</td>
<td>0.12</td>
<td>0.18</td>
<td>0.23*</td>
<td>0.45*</td>
</tr>
<tr>
<td>USJCE28</td>
<td>0.16</td>
<td>0.25*</td>
<td>0.32*</td>
<td>0.49*</td>
</tr>
<tr>
<td>USJCS28</td>
<td>0.17*</td>
<td>0.30*</td>
<td>0.36*</td>
<td>0.48*</td>
</tr>
<tr>
<td>USJCPD28</td>
<td>0.16</td>
<td>0.25*</td>
<td>0.29*</td>
<td>0.50*</td>
</tr>
<tr>
<td>USJIE</td>
<td>0.12</td>
<td>0.23*</td>
<td>0.31*</td>
<td>0.51*</td>
</tr>
<tr>
<td>USJS</td>
<td>0.15</td>
<td>0.27*</td>
<td>0.32*</td>
<td>0.51*</td>
</tr>
<tr>
<td>USJP</td>
<td>0.11</td>
<td>0.20</td>
<td>0.24</td>
<td>0.44*</td>
</tr>
</tbody>
</table>

*p < 0.01; *p < 0.05.

HAQ, Health Assessment Questionnaire; VASP, visual analogue scale for pain; VASOA, visual analogue scale for patient’s overall assessment; ESR, erythrocyte sedimentation rate; CRP, C reactive protein; TJC, tender joint count; SJC, swollen joint count; SJI, swollen joint index; USJCE, ultrasonographic joint count for effusion; USJCS, ultrasonographic joint count for synovitis; USJCPD, ultrasonographic joint count for power Doppler signal; USJIE, ultrasonographic joint index for effusion; USJSI, ultrasonographic joint index for synovitis; USJIPD, ultrasonographic joint index for power Doppler signal.
archiving computer system. The saved images were blindly read by the same rheumatologist who performed US examination (EN) 3 months after the initial scanning, and by a fourth rheumatologist (JU) trained in US. These images did not show measurements. Before the study the investigators reached a consensus about the US scales. Figure 1 shows some examples of the US findings.

Statistical analysis

A $\chi^2$ test was applied for comparing cumulative variables, and Student’s $t$ test and Pearson’s correlation were used for continuous variables. Any value of $p < 0.05$ was considered significant.

Interobserver agreement between the clinical investigators and the US investigators and intraobserver US agreement were calculated using an unweighted $k$ test. The $k$ value measures agreement between pairs of observers, eliminating random concordance. A $k$ value $< 0.40$ was considered poor, 0.40–0.60 moderate, 0.60–0.80 good, and 0.80–1 excellent.

RESULTS

Patients’ characteristics

Rheumatoid factor was positive in 73/94 (78%) patients. Therapeutic regimens included non-steroidal anti-inflammatory drugs in 60 (64%) patients, corticosteroids in 56 (60%), methotrexate in 61 (65%), lefunomide in 18 (19%), sulfasalazine in 10 (11%), antimalarial drugs in 10 (11%), azathioprine in 2 (2%), and ciclosporin in 2 (2%).

Table 5 shows the correlations between clinical, laboratory, and US variables. TJC, SJC, and SJI correlated poorly with HAQ, VASP, and VASOA ($p < 0.01$). TJC did not correlate with ESR or CRP ($p < 0.04$), nor did SJC and SJI correlate poorly with ESR and moderately with CRP ($p < 0.01$).

No significant differences were found between patients examined with scanners 1 and 2 for mean age, mean duration of symptoms, mean VASP and VASOA scores, mean joint count for tenderness and swelling, mean SJI, mean HAQ score and mean ESR and CRP values (table 2).

All joints, including nine prosthetic joints, could be easily assessed by US. The US examination took 20–30 minutes for each patient, not including documentation.

Clinical and ultrasonographic joint involvement

US detected more joints with effusion and synovitis than clinical examination. Mean (SD) USJCE was 15.2 (9.3), mean USJCS 14.6 (9.4), and mean SJCS 11.5 (7.4) ($p = 0.003$ for mean USJCE vs mean SJCS and $p = 0.01$ for mean USJCS vs mean SJCS, respectively).

Table 4 shows the correlation between clinical, ultrasonographic, and laboratory variables as measured by US.

No significant differences were found between patients examined with scanners 1 and 2 for mean age, mean duration of symptoms, mean VASP and VASOA scores, mean joint count for tenderness and swelling, mean SJI, mean HAQ score and mean ESR and CRP values (table 2).

All joints, including nine prosthetic joints, could be easily assessed by US. The US examination took 20–30 minutes for each patient, not including documentation.
TJC and SJC correlated weakly (r = 0.49, p<0.01). TJC did not correlate with joint count for US effusion, US synovitis, and PD signal (p>0.05). SJC correlated with US joint counts (r = 0.58, 0.57, and 0.56, respectively, p<0.01). Correlations between SJ and US index for effusion, synovitis, and PD signal were better (r = 0.65, 0.64, and 0.58, respectively, p<0.01). A high correlation (r >0.80, p<0.01) was found between all the US variables.

TJC and SJC correlated highly with TJC28 (r = 0.95, p<0.01) and SJC28 (r = 0.96, p<0.01). In the same way, there was a high correlation between USJCE and USJCE28 (r = 0.92, p<0.01), USJCS and USJCS28 (r = 0.92, p<0.01), and between USJCPD and USJCPD28 (r = 0.94, p<0.01), USJCE28, USJCS28, and USJCPD28 showed similar correlations with clinical and laboratory measures as with the corresponding 60 joint counts (table 3).

For the 69 patients examined with scanner 1, US findings showed a similar correlation with clinical and laboratory measures as those obtained in the overall group (table 4). There was a high correlation between US parameters and CRP levels (p<0.01).

Clinical and ultrasonographic reliability
Clinical interobserver agreement ranged from poor to excellent (table 5). Overall interobserver agreement for tenderness was better than for swelling and for the swelling index. There was a high level of intraobserver and interobserver agreement, from moderate to excellent, for US effusion, synovitis, and PD signal (table 6).

DISCUSSION
We found that US was clearly more sensitive than physical examination in detecting joint swelling. In addition, US findings correlated better with CRP and ESR than clinical joint swelling. In keeping with our results, other studies have demonstrated that US detects subclinical synovitis. Because therapeutic decisions depend considerably on clinical synovitis, the undetectable synovitis may explain the continued bone damage found in patients with clinically controlled RA. In the same way, the results of Jevtic et al confirmed that in joints with inflammatory active pannus detected by contrast enhanced MRI, progression of bone-destructive changes is expected.

Hypervascularisation and angiogenesis of the synovial membrane are considered to be primary pathogenic mechanisms responsible for invasive and joint destructive behaviour of rheumatoid pannus. Dynamic contrast enhanced MRI findings have demonstrated a close correlation with histological signs of knee synovial inflammation in patients with RA. However MRI is expensive, time consuming, and not widely available for routine use in most countries. Recently, PD US has demonstrated a high sensitivity (88.8%) and specificity (97.9%) for the assessment of inflammatory activity in the MCP joints of patients with RA compared with dynamic contrast enhanced MRI. One advantage of US over MRI is that examination of all peripheral joints can be done as many times as required and prosthesis or implants do not interfere with US images. Last but not least, rheumatologists can be trained to perform US, removing the need for referral to a radiologist and thus saving time and money.

US variables correlated with clinical joint swelling, CRP, and ESR, correlated weakly with VAS scales, and did not correlate with clinical joint tenderness and HAQ. Previous studies comparing clinical and US assessment have also reported a stronger correlation between US and physical examination of joint swelling than between US findings and patient’s perception of joint tenderness. Furthermore, Qvistgaard et al found that the degree of synovial vascularisation in finger joints detected by colour Doppler correlated with ESR, and not with VAS for pain, VAS for patient overall assessment, and HAQ scores in patients with RA. In fact, joint tenderness, VAS scales, and HAQ scores indicate either disease activity or structural joint damage or deformity secondary to previous synovial inflammation not present when clinical evaluation is performed.

This study evaluated 60 joints in each patient with RA, whereas similar previous reports investigated only a small number of joints, such as knee, MCP, and PIP joints. We considered that 60 joint counts represented an overall assessment of disease inflammatory activity. However, reduced joint counts for tenderness and swelling, such as the 28 joint count are widely used in the evaluation of RA inflammatory activity in daily practice and in clinical trials. Our 28 joint count for US effusion, synovitis, and PD signal correlated well with the 60 joint count. Thus, the reduced joint count could be used in US evaluation in rheumatological practice. Although effusion and synovitis can also be seen in other periarticular locations of the examined joints, we performed a simplified US investigation in order to shorten scanning duration. The proposed US evaluation of 28 joints can be performed in 15 minutes in daily practice.

Recent studies have reported the validity and reliability of cheaper US machines for assessing rheumatoid synovial inflammation, though most reports have used expensive machines like our scanner. Analysis of the results from scanner 1 showed that they were comparable with the overall
results. Therefore, a considerably more affordable machine like scanner 1 could also be used.

Some limitations of our study should be mentioned. The assessment of a single selected US image instead of a real time examination of the joints performed by the second rheumatologist obviously introduces bias into the study. However, this is the standard way to record US examination in daily practice and the images for a second reading were chosen by an experienced sonographer. Our data are in accordance with those of Szukladerek et al, who recently reported a high interobserver agreement for the identification and semiquantitative measurement of effusion, synovitis, and PD signal in the small joints of patients with RA. In addition, the rheumatologist obtaining the sonograms could not be completely unaware of a patient’s joint signs and symptoms. To avoid as much bias as possible, the US examination was carried out in the dark.

A number of factors potentially limit the use of PD. Firstly, no examination protocols or standard settings for PD US machines exist. In addition, PD US is extremely sensitive to tissue movement, especially at low PRF, which can result in “flash” artefacts. However, we used pulsed Doppler spectra as proof of the presence of vessels when images were doubtful.

CONCLUSION

We propose that the combination of grey scale US and PD could be used as a sensitive and reliable non-invasive and widely available method complementary to standard clinical assessment for evaluating rheumatoid synovial inflammation in daily management and clinical trials. Moreover, images of the examinations performed can be kept. PD may become a cost effective alternative to gadolinium enhanced MRI.

Longitudinal studies that correlate PD findings with long term clinical changes and radiographic erosive joint damage in patients with RA are highly warranted.

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