Doppler ultrasound (DUS) is increasingly used in the evaluation of joint inflammation and has the advantage of giving an objective analysis of the disease without the use of ionising radiation. It is accessible in many outpatient clinics, has no contraindications, and poses no problems regarding patient compliance. Colour and power Doppler have been used to identify hyperaemia associated with inflammation by visualising the vascularity in the inflamed synovial membrane. DUS can detect synovitis in arthritis, and several studies in patients with rheumatoid arthritis (RA) have shown that it can detect treatment response following glucocorticosteroid administration and after systemic treatment with biologicals such as tumour necrosis factor-α. Because the newer US machines are equipped with high frequency transducers (10–20 MHz) and increased Doppler sensitivity, it has become possible to detect vascularity even in a normal synovial membrane, as previously shown in the elbow joint. Both the detection limits and the reference range for DUS in normal synovial perfusion have yet to be established.

As the wrist and fingers are frequently targeted in RA, the aim of this study was to examine the wrist and finger joints of healthy volunteers by DUS for presence of colour pixels on the scan (indicating perfusion) and to evaluate the flow profile of vessels if detected, in the normal synovial membrane.

MATERIALS AND METHODS

Subjects
Staff members in our clinic and their families made up the 28 subjects we recruited to represent predefined age groups of 10 year intervals from 15–65 years or above. A total of 27 healthy volunteers (15 women, 12 men; mean age 45 years, range 18–93) were included in the study. The subjects were not currently engaged in heavy manual labour or performing sports activities with their hands, had no history of arthritis or hand or wrist trauma, and were not experiencing current symptoms. None of the subjects had clinical signs of inflammatory or degenerative joint diseases. A total of 324 joints were examined (12 joints per subject): the wrist, carpometacarpal 1 (CMC1), metacarpophalangeal (MCP), and proximal interphalangeal (PIP) joints of healthy controls.

Methods: Twenty seven healthy volunteers (15 women, 12 men; mean age 45 years, range 18–93) with a total of 324 joints were examined by DUS. The synovial vascularity was determined by colour Doppler and the spectral Doppler resistive index (RI). Patients were only included if no synovitis was suspected at the clinical examination and the values for biochemical analysis were within the normal range.

Results: We found colour pixels in 15 of the 27 examined wrist scans and in 8 of these a measurable RI. In the CMC1 joint, colour pixels were found in 11 of the 27 scans and in 9 of these a measurable RI. For the MCP joints, colour pixels were found in 10 out of 135 scans and in 3 of these a measurable RI. The mean RI for all three types of joints was 0.85 and the mean pixel fraction varied from 0.05 to 0.08. Only one PIP joint scan was found to have colour pixels and a flow with an RI of 0.67.

Conclusion: Synovial vascularity may be detected in healthy subjects using DUS. Newer US machines have higher Doppler sensitivity, and it is necessary to be able to distinguish normal from pathological synovial flow.
were scanned. The finger joints were scanned in the longitudinal plane only and the palmar aspects were not investigated. The CMC1, MCP1, and PIP joints were scanned in an arch of 180° from the ulnar to the radial side. MCP2–5 were scanned in the regions that were accessible from the dorsal side: MCP2 and MCP5 in an arch of 150° and MCP 3–4 in an arch of 120°.

In this study, synovial hypertrophy or the synovial volume was not estimated as previously described, as the aim was to obtain the Doppler images only. No normal values so far exist for the thickness of the normal synovial membrane, and it is difficult to differentiate between cartilage and synovial membrane both in patients with arthritis and in normal joints. The cartilage is therefore included in the joint volume and we see this as a systematic error. However, as there is no vascularisation of the cartilage it does not have any influence on the Doppler findings.

**Colour Doppler**

Blood flow in the synovial membrane was visualised with colour Doppler, which is as sensitive as the power Doppler on the Sequoia®. The colour Doppler settings were the same for all joints and all volunteers, with a gain setting just below the noise level using our settings for low flow (Nyquist limit ± 0.014 m/s and 7 MHz Doppler frequency). With these settings, all the colour pixels in the image correspond to motion—that is, blood flow. That the colour pixels represented flow and not an artefact was established with the spectral Doppler displaying arterial or venous flow or, in the few cases where the signal was too weak for spectral analysis, by the constant position of the colour pixels inside the synovial membrane. This in contrast to noise or movement artefacts that appear in random fashion and in clear contrast to artefacts generated at highly reflective surfaces.

Quantitative estimation of the vascularisation in the synovial membrane was performed using the selected colour Doppler image (CDI) with maximum colour activity. The digitally stored CDI in DICOM format was transferred to a processing program (a beta version of DataProTM; Noesis, Courtabeuf, France). The region of interest (ROI) was traced on the synovial membrane. On the US image the synovial membrane appears predominantly hypoechoic, covering the bone surface adjacent to the joint and superficially bounded from the subcutaneous tissue by a thin white strand, the fibrous capsule. The US joint anatomy is illustrated in fig 1. The number of colour pixels and the total number of pixels in the ROI were counted, and the number of colour pixels was then expressed in relation to the total number of pixels in the ROI as the colour fraction.

**SPECTRAL DOPPLER**

Previous work with both the pulsatility and resistive indices have lead us to work only with the resistive index (RI) as it is easy to understand.

Using colour Doppler as guidance, the spectral Doppler sampling area was placed over an intra-synovial artery and the spectrum was obtained. An intra-muscular artery in the thenar muscles was sampled as a reference. The US unit then calculated the RI as (peak systolic flow minus end diastolic flow)/peak systolic flow. If more than one synovial arteries could be sampled, a mean RI was used. With the spectral Doppler technique it is possible to evaluate the type of flow—that is, low peripheral resistance (low peripheral resistance, which is normal in resting musculoskeletal tissues). Because the intra-synovial vessels are very small, both the artery and its concomitant veins are often sampled simultaneously even with the smallest possible Doppler gate. A flow reversal during diastole (normal in musculoskeletal tissues) will then go unnoticed because the reversed arterial flow will drown in the venous signal. In order to obtain uniform measurements, we therefore limited the spectral measurements to the arterial side of the Doppler line and defined 1.00 as the maximum for RI.

When spectral Doppler measurements could not be measured, the RI was noted to be the same as in the thenar muscles, as the resistance in the synovial arteries was presumed to be the same as extra-synovial musculoskeletal flow. This assumption has been made for statistical reasons to obtain a value for normal RI in the cases where no flow was present on colour Doppler. No flow on colour Doppler is a normal finding and for RI we therefore used a value obtained from normal resting musculoskeletal tissue—that is, the thenar muscles in this study.

Erosions visualised by US were defined as a disruption of the bone surface by an indentation, which gave the bone surface an irregular appearance. Osteophytes were defined as bone extensions from non-cartilaginous bone surfaces inside the joint space.

**STATISTICS**

Statistical analysis was performed using the SPSS program (version 10.0). For correlation between age and US parameters and between the two US parameters, Pearson’s test was used. Student’s t test was used for evaluation of the total dataset. Level of significance was p = 0.05.

**RESULTS**

There were no differences between the US findings in healthy men and women. The DUS findings for the individual joints are shown in table 1. The mean RI and the colour fraction showed a statistically significant correlation, with R values of −0.518 (p<0.01).

<table>
<thead>
<tr>
<th>Joint</th>
<th>Mean RI (range)</th>
<th>Colour Fraction (range)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMC1</td>
<td>0.50 (0.16)</td>
<td>0.00 (0.00)</td>
</tr>
<tr>
<td>MCP1</td>
<td>0.60 (0.28)</td>
<td>0.01 (0.00)</td>
</tr>
<tr>
<td>PIP2–5</td>
<td>0.60 (0.28)</td>
<td>0.01 (0.00)</td>
</tr>
</tbody>
</table>

Mean (SD) (range) RI for all joints for the healthy volunteers was 1.00 (0.06) (0.47–1.00) and the colour fraction 0.008 (0.03) (0–0.32). The mean (range) RI from the thenar muscles was 1.00 (1.00–1.00).

Colour Doppler activity was detected in 11% (37/324) of the joints (fig 2). Of these, it was possible to obtain spectral Doppler signals in 21 joints. Mostly, only one solitary artery could be sampled and in only three joints (two wrists and one CMC1 joint) could three arteries be sampled.

There was a weak, but statistically significant correlation between age and colour fraction (R value 0.163; p<0.01), but this correlation was not significant if the oldest subject was excluded. We found no correlation between age and RI.

Small erosions were found in three CMC1 joints but not in any other joints and in no particular age group. In these three joints there was colour Doppler activity in one (with the highest colour fraction (0.32)) and three arteries detected with spectral Doppler (mean RI 0.57). OA changes (osteophytes) were found in two CMC1 joints and in 12 PIP joints in seven subjects, all >52 years of age. The two CMC1 joints had colour Doppler activity with colour fractions of 0.01 and 0.02. Both had measurable flow with RI of 1.00 and 0.71. None of these OA joints had erosions.

**DISCUSSION**

The present study demonstrated a presence of Doppler activity in 11% of the hand and finger joints of healthy
volunteers with no history of arthritis, hand or wrist trauma, or current symptoms. Although the majority of the joints examined were without Doppler activity, we believe the 11% positive in this study to be substantial, disqualifying the mere presence of colour pixels on scans as an indication of inflammation.

Anatomically, the synovial membrane consists of a lining layer and a sub-lining layer bordering on the fibrous capsule. Deep in the sub-lining, arteries and veins can be seen, which decrease in size as the lining layer is approached. Detection of arterial flow in the synovial membrane depends on the sensitivity of the Doppler equipment, and our findings of colour pixels in normal synovial membrane scans are in accordance with one previous study reporting colour pixels in joint scans of 15 medical students, but in contrast to other studies reporting negative Doppler findings in healthy controls.

Doppler US is a relatively new tool in the investigation of local inflammation in arthritis. The Doppler signals are a reflection of the number of erythrocytes that move within the image plane and the speed with which they move. In consequence, the colour pixels are an estimate of the amount of moving blood in the investigated area.

Previous studies have shown that there is a correlation between the histological and sonographical degree of vascularity; however the number of colour pixels do not correspond to the number of vessels visualised by DUS but rather to the perfusion of these vessels. A correlation between US and MRI has also been determined recently, and the inflammatory parameters used in this study has been validated in a recent study comparing US and MRI, in which both symptomatic and asymptomatic joints were examined.

The presence of colour pixels on scans of the synovial membrane has until now been taken as a sign of hyperaemia, but with increasing Doppler sensitivity, normal synovial flow may also result in colour pixels. Accordingly, the mere presence of colour pixels cannot at present be interpreted as a sign of synovitis per se, especially in the wrists and to some extent in the MCP joints as indicated by our study. The number of colour pixels seen in our scans is comparable to that which we frequently encounter sporadically when we scan the asymptomatic joints of the hands of an RA subject. We do not consider it a sign of pathology, and as we show here, these findings can also be seen in healthy volunteers. Spectral Doppler examinations on joints of healthy volunteers have, to our knowledge, not been previously investigated. The spectral Doppler examination depends on colour Doppler to identify the presence and location of vessels and was therefore only applied in the 37 colour Doppler positive joints in this study. In 21 of these
joints, it was possible to identify an artery and thereby obtain an RI.

In normal resting musculoskeletal tissues, flow is seen with colour or power Doppler in muscles and connective tissue, seldom in joints, and never in tendons. When the flow is assessed by spectral Doppler, no diastolic flow is found in these tissues at rest and consequently, the RI is 1.00. This was demonstrated in this study by all 27 subjects having an RI of 1.00 in the thenar muscles. In rheumatology, low values of RI indicate pathology, which has been observed in previous studies. Among the healthy volunteers only one synovial artery could be sampled from the investigated joints, apart from two wrists and one CMC1 joint in which three synovial arteries could be sampled. In general, when examining RA patients with our method, inflamed joints have three or more different synovial arteries that can be sampled. The present findings show that even though low RI values may be obtained from healthy volunteers, our mean values (0.85) are higher than those reported in RA patients (0.4–0.7) and 0.65–0.76.

The positive Doppler findings in some normal joints indicate that the mere presence of colour pixels on scans may not be enough for diagnosis of inflammation. In chronic synovitis we also find greyscale abnormalities such as synovial proliferation and/or effusion, but if DUS is to be used in the detection of inflammatory changes in very early arthritis it is not likely in those situations to find synovial proliferation and/or effusion. Therefore, it seems rational that a cutoff level (above which inflammation is present) for both colour fraction and RI should be established for diagnostic purposes rather than depending on the more or less static greyscale changes.

In these asymptomatic subjects, we found few greyscale changes indicating OA. These changes were predominantly in PIP joints and not accompanied by colour Doppler activity. In the two CMC1 joints with osteophytes we did find colour Doppler activity (colour fraction of 0.01 and 0.02) and in one joint a low RI (0.71) also. All these joints were identified in subjects over 50 years of age. The only signs of erosive changes in healthy volunteers were found in the CMC1 joint (three cases), which is also prone to OA changes. In the joints recommended by the ACR to be evaluated in RA we found no erosive changes, indicating that the presence of erosions by US in the wrist, MCP, and PIP joints are signs of pathology.

Our findings show that colour Doppler activity is most common in the wrist (15/27) followed by CMC1 (11/27), MCP (10/135), and very seldom in the PIP joints (1/135). The rather frequent observation of positive findings in the wrist may be due to the composite structure of this joint. If it is assumed that joints of comparable size have the same chance of displaying colour Doppler activity, the wrist is naturally more likely because it is a group of joints. In this respect, CMC1 has remarkably high colour Doppler activity, which, according to joint size, should be closer to the MCP joints. In this study, the CMC1 may therefore be seen as an outlier not only in terms of colour Doppler activity but also in OA changes and erosions. In accordance with the ACR recommendations from the clinical point of view, the ultrasonographic changes are in support of an exclusion of this joint from a diagnostic evaluation of arthritis by imaging.

CONCLUSION

Colour Doppler activity may be found in healthy volunteers, most commonly in the wrist, more rarely in the MCP joints, and hardly ever in the PIP joints. The CMC1 joint is prone to both greyscale and Doppler findings not necessarily related to RA and therefore it is reasonable that this joint should not be
part of the recommended joint evaluation. The Doppler sensitivity is increased in newer US machines and seems to approach the detection limit of normal synovial flow. As sensitivity is likely to continue with development of new machines, it will be necessary in the near future to be able to distinguish normal from pathological synovial flow.

Authors’ affiliations
L Terslev, S Torp-Pedersen, E Qvistgaard, P von der Recke, H Bliddal, The Parker Institute, Department of Rheumatology, Frederiksberg Hospital, DK 2000 Frederiksberg, Denmark

REFERENCES