Knee joint synovitis: study of correlations and diagnostic performances of ultrasonography compared with histopathology

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ABSTRACT

Objectives Ultrasonography (US) is a fast, available and low-cost imaging tool used for detecting knee synovitis. Our aims were to assess the relationship between US and histology findings in appraising levels of inflammation and vascularity in the knee joint in subjects with inflammatory arthropathies; to determine whether differences exist in the appraisal between varying knee compartments and to compare US performances compared with gold standard histology for knee synovitis detection.

Methods Subjects with actively inflamed knee joint having crystal arthropathies, rheumatoid arthritis, psoriatic arthritis or osteoarthritis were prospectively recruited from rheumatology clinics after giving their written consent between May and October 2015. Study was approved by the institutional ethics committee. The knee was divided into three compartments (medial, lateral, superior). Patients had a knee US followed by a knee arthroscopy with biopsies retrieval from each compartment. Biopsies were blindly scored for lining layer hyperplasia, inflammation, vascularity, CD68 and factor VIII staining. Correlation was determined using the Spearman’s correlation test.

Results 26 patients with active arthritis in a knee joint and recent onset of disease were prospectively included. Strong correlations were observed between US synovitis grade and histological inflammation score (r=0.63; P=0.002), US Doppler grade and histological score for vascularity (r=0.68; P<0.001); US measured synovial thickness and lining layer hyperplasia (r=0.61; P=0.002). Moderate correlation was found between US synovitis grade and CD68 score (r=0.49; P=0.02).

Conclusion US findings correlate with histological inflammation and vascularity scores in actively inflamed knee joints. US accurately describes knee synovitis.

INTRODUCTION

Quantifying inflammation, disease activity and predicting joint destruction in rheumatic diseases remains a significant challenge.

Clinical examination remains important, and although this can contribute in assessing synovitis and predicting outcome, it remains insensitive for both these parameters.1

Ultrasonography (US) allows detection of active synovitis using B Mode for semi-quantitative synovitis grade assignment and Doppler mode for hypervascularisation detection.2–4 Therefore, US examination allows the evaluation of disease activity5 and the determination of treatment efficacy.6 7 Some studies have assessed the reliability of US in appraising levels of joint inflammation. Karim et al have demonstrated that US examination is reproducible and has a better sensitivity (98%), specificity (88%), positive (98%) and negative (88%) predictive values than clinical examination, using arthroscopy as the gold standard.9 Other studies showed a better reliability of US examination compared with clinical assessment for synovitis detection in patients with rheumatoid arthritis (RA).9 10 MRI assessment of synovitis has been shown to correlate with histological inflammation in synovitis.11 MRI performances in prediction of both further diagnosis of RA in patients with undifferentiated arthritis (UA) and structural

Key messages

What is already known about this subject?

► Correlations between US findings and some histological findings exist in small joints.

What does this study add?

► Correlations exist between US findings and histological findings such as inflammation score and CD68 staining, in knee joints.

How might this impact on clinical practice?

► Ultrasonography in large joints such as knee is a reliable tool.

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prognosis in established patients with RA have been demonstrated.\textsuperscript{12–14}

Histological features such as levels of macrophage infiltration and lining layer hyperplasia are associated with disease activity and structural damages.\textsuperscript{15} Validated scores exist in order to assess and quantify histological inflammation within the synovial membrane.\textsuperscript{16,17}

Disease activity score 28 has been showed to correlate with histological inflammation of the synovial membrane. Indeed, sublining macrophages are considered a sensitive biomarker of response to treatment.\textsuperscript{18}

However, synovial biopsies are not routinely performed in order to assess inflammation. In contrast, non-invasive imaging tools are routinely used.

US is widely performed and recognised as a reliable assessment tool for synovitis in rheumatology. However, only a few studies have assessed the correlation between Doppler mode features and histological vascularisation, and mostly in long-lasting RA populations.\textsuperscript{19–22} Moreover, it remains to be clarified whether US findings are a helpful biomarker for disease activity and possibly treatment response compared with synovial histology.

Our prospective study aimed to assess the correlation between US and histology findings in appraising levels of inflammation and vascularity in different areas of the knee joint in subjects with inflammatory arthropathies; to determine whether differences exist in the appraisal between varying knee compartments and to assess diagnostic performance of US measures for diagnosis of synovial inflammation compared with the histological gold standard.

METHODS

Patients and methods

Patients with an actively inflamed knee joint and recent onset of disease were consecutively recruited from the clinics of the Rheumatology Department, St. Vincent's University Hospital. All patients gave fully informed written consent for US evaluation and synovial biopsy by needle arthroscopy of the knee for research purposes as approved by the St. Vincent's Healthcare Group Medical Research and Ethics Committee.

All patients referred to outpatient clinics with an actively inflamed knee joint, having both UA or recently diagnosed with osteoarthritis (OA), autoimmune inflammatory arthropathy or crystal arthropathy were prospectively included from May to October 2015. All patients had an US examination of the knee joint immediately before undergoing arthroscopy of the same knee. The knee was divided into three compartments. The three compartments were defined as follows:\textsuperscript{8} medial compartment (MC) was defined as ‘the area medial to the patella running inferiorly past the medial joint line to the inferomedial aspect of the joint capsule’. The lateral compartment (LC) was defined as ‘the area lateral to the patella running inferiorly past the lateral joint line to the inferolateral aspect of the joint capsule’. The superior compartment (SC) was defined as ‘the area above the patella’ (figure 1).

US evaluation

One trained physician (3 years’ experience) performed the US examination blindly from clinical data using a General Electrics Logiq 9 US machine with a 8–12 MHz linear transducer. US evaluation was performed on the knee in 20°–30° flexion according to the recommendations of the EULAR Musculoskeletal Ultrasound working group. A pulse repetition frequency of 750 Hz was used for power Doppler examination. Power Doppler signal (PDS) gain was set manually to a maximum of sensitivity just below the disappearance of colour noise.

Each compartment was scanned in longitudinal and lateral planes.\textsuperscript{2} A well-established definition for joint effusion and synovial hypertrophy was used.\textsuperscript{23} The joint effusion evaluation was binary (present or not present). A cut-off limit of a longitudinal scan measure >4 mm was used for pathological effusion definition.\textsuperscript{24} The evaluation for synovitis was performed using well-established semiquantitative scores with a four-grade scale of 0–3 for both B mode and Doppler examination\textsuperscript{9} for each of the three compartments. Synovial thickness was measured with longitudinal scan for SC and transversal scans for MC and LC (figure 2).

B mode 0–3 grading for synovitis was 0=absence of synovial thickening, 1=mild synovial thickening, 2=moderate synovial thickening and 3=marked synovial thickening. Power Doppler activity was graded from 0 to 3 as follows: 0=absence of signal, no intra-articular flow;
Imaging

1=mild (2–3 signals, including two confluent vessels); 2=moderate confluent vessels in <50% of the synovium; and 3=marked vessel signals in >50% of the synovium.9 10 The median of the scores attributed to each compartment was then calculated as an overall score for each patient.

Arthroscopy
Arthroscopy was performed by two trained experienced operators who were blinded from US findings, using a Wolf (Germany) or a Storz (Tuttlingen, Germany) 2.7 mm rigid arthroscope as previously described.24 The procedure was performed under local anaesthetic. Each compartment of the knee was marked according to previous definition before the procedure. Position of the arthroscope and forceps within the targeted compartment was ensured by transilluminating the skin with the arthroscope and impressing move under skin with forceps.

A 100 mm visual analogue scale (VAS) was used to score each compartment for both synovitis and vascularity.8 An average of 20 biopsies were retrieved from each compartment and labelled. The procedure was entirely recorded and stored for each patient.

Biopsies processing and staining
Tissue was fixed in formalin, embedded in paraffin and 3 μm sectioned (Microtome). Sections were baked for 30 min at 90°C, deparaffinised in xylene and dehydrated in alcohol and distilled water. Sections were then incubated with primary antibodies against human CD68 or human FVIII (Dako, Denmark) for 2 hours. Slides were then incubated with horseradish peroxidase-conjugated secondary antibody (Dako) for 30 min. Diaminobenzidine tetrahydrochloride was used as the chromogene substrate (Dako). Slides were counterstained with haematoxylin (BDH Laboratories, Poole, UK) and mounted. Each slide was then scored by one trained experienced operator, blindly from US and arthroscopic findings, for inflammation and vascularisation using a semiquantitative five-scale score (0=Minimal infiltration, 1=Mild infiltration, 2=Moderate infiltration, 3=Severe infiltration, 4=Infiltration with numerous inflammatory cells).25–28 The lining layer hyperplasia was assessed by calculating the mean number of cell layers. CD68 slides were read and scored using a semiquantitative five-scale score for both lining layer and sublining calibrated against a standardised atlas.25 26 28 The FVIII stained slides were scored by counting the absolute number of blood vessels per high-powered field (magnification ×20) expressing FVIII. The score was the mean number calculated of five fields randomly selected29 (figure 2).

Statistical analyses
Normally distributed continuous data were expressed as the mean with SD or for non-normally distributed data as median with IQR. Professional statistician was consulted for statistical design. Correlation were studied using the Spearman’s correlation tests. When the correlation coefficient was r<0.3, correlation was considered as mild. When comprised between 0.3<r<0.6, the correlation strength was considered as moderate. When the correlation coefficient was comprised between r>0.6, the correlation strength was considered as high. A P value <0.05 was considered statistically significant. Correlation analyses were performed through GraphPad Prism V.6.0. Sensitivity, specificity, predictive positive and negative values of US to detect inflammatory arthritis compared

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Figure 2  Example of ultrasound, histological and CD68 or factor VIII (FVIII) analyses. (A) B mode synovial thickness measurements. (B) H&E stained slide. Histological synovitis: lining layer hyperplasia, stroma activation and inflammatory infiltrate. (C) Corresponding CD68 staining. Original magnification 20×. (D) Power Doppler activity assessment. (E) H&E stained slide. Multiple blood vessels. (F) Corresponding FVIII staining. Original magnification 20×.
with histological gold standard were calculated through SPSS software.

**RESULTS**

**Patients' characteristics**

Twenty-six patients were included between May and October 2015 in Saint Vincent's University Hospital. Demographic and clinical characteristics of the patients are shown in table 1.

**US, arthroscopic and histological scores**

Median US synovitis grade was 1 (0–2). Median measured synovial thickness was 1.6 mm (1.1–3.2). Median histological scores and CD68 and FVIII scores are shown in table 2.

**Correlation between US and histological scores**

**US and histology**

In the LCs, we found strong correlations between US B mode synovitis score and histological inflammation score (r=0.67; P=0.013), and between US measured synovial thickness and histological score for lining layer hyperplasia (r=0.55; P=0.038).

In the SC and MC, correlations between US B mode synovitis score and histological inflammation score, and between US measured synovial thickness and histological score for lining layer hyperplasia did not reach statistical significance (table 3).

Median scores represent the median of for lateral, medial and superior scores.

Strong correlations were observed between median US B mode synovitis score and median histological inflammation score (r=0.63; P=0.001); median US measured synovial thickness and median histological lining layer hyperplasia score (r=0.61; P=0.002); and median US power Doppler score and median histological vascularisation score (r=0.68; P=0.0003) (table 3).

| Table 1 | Patients' characteristics |
|---------|---------------------------|
| **Total** | 26                        |
| **Demographics** |                       |
| Gender |                           |
| Female | 21 (80.8%)                |
| Mean age (years) (range) | 50.8 (31.3–68.2)        |
| Female mean age (years) (range) | 50.8 (31.3–68.2) |
| Male mean age (years) (range) | 50.7 (35–57.9)         |
| Biopsied knee |                    |
| Right | 13 (50.0%)                |
| **Diagnosis** |                  |
| RA | 17 (65.4%)                |
| Rheumatoid factor positive | 10 (38.6%)           |
| Anti-CCP positive | 12 (46.2%)            |
| DAS 28 (mean) (SD) | 3.85 (1.03)            |
| PsA | 3 (11.5%)                 |
| OA | 4 (15.4%)                 |
| Gout | 1 (3.8%)                 |
| UA | 1 (3.8%)                  |
| Elevated CRP (> 5 mg/L) | 14 (53.8%)            |
| Erosive disease (RA) | 4 (15.4%)             |
| Treatment (RA, PsA) |                   |
| Methotrexate | 3 (11.5%)            |
| TNF inhibitors | 4 (15.4%)             |
| Other | 1 (3.8%)                  |
| CCP, cyclic citrullinated peptides; CRP, C reactive protein; DAS, disease activity score; OA, osteoarthrosis; PsA, psoriatic arthritis; RA, rheumatoid arthritis; TNF, tumour necrosis factor. |

| Table 2 | Ultrasonography (US) and histological findings |
|---------|---------------------------------------------|
| Median US synovitis grade (0–3 semiquantitative score) (Q1–Q3) | 1 (0–2) | 1 (1–2) | 1 (0–2) | 1 (0–2) |
| Median measured synovial thickness (mm) (Q1–Q3) | 1.55 (0.8–4.3) | 2.1 (1.3–3.1) | 2.05 (0.3–3.4) | 1.6 (1.1–3.2) |
| Median US synovitis grade (Doppler mode) (0–3 semiquantitative score) (Q1–Q3) | 0 (0–2) | 0 (0–1) | 0 (0–2) | 0 (0–2) |
| Median histological synovitis score (0–4) (Q1–Q3) | 2 (1–2) | 1 (1–2.8) | 2 (1–3) | 2 (1–2.5) |
| Median CD68 score (Q1–Q3) | 3 (2.75–4) | 3 (2–3) | 3 (2–3) | 3 (2–3.5) |
| Mean number of vessels per slide (absolute number with FVIII staining) (SD) | 8.3 (5.4) | 11 (6.3) | 16 (8.0) | 10.2 (6.3) |

*Overall scores are the mean scores of the three compartments. FVIII, factor VIII staining; Q1–Q3, first and third quartiles.
US and immunohistochemistry
In the SC, LC and MC, correlations between US B mode synovitis score or US measured synovial thickness and CD68 staining did not reach statistical significance (table 3).

However, moderate correlation was observed between median US B mode synovitis score and median score for CD68 staining \((r=0.49; P=0.02)\); strong correlations were observed between median US measured synovial thickness and median score for CD68 staining \((r=0.59; P=0.006)\) (table 3).

In the LC, significant strong correlation was found between US power Doppler score and FVIII staining \((r=0.74; P=0.009)\). However, median US power Doppler score and median FVIII staining did not reach statistical significance (table 3).

Correlation between the three knee compartments for US, arthroscopic and histological findings
Strong correlations were observed between the SC, MC and LC of the knee for US synovitis grade and US measured synovial thickness (table 4). However, power Doppler grade partially correlates only (table 4).

Table 3 Correlation between ultrasonography and histological findings

| Compartment | Histological inflammation score | Histological lining layer thickness score | Histological vascularisation score | CD68 score | FVIII score | Power Doppler grade |
|-------------|---------------------------------|------------------------------------------|----------------------------------|------------|------------|--------------------|
| Superior    | 0.48†                           | 0.35                                     | 0.27                             | 0.42†      | 0.37       | 0.20               |
| Medial      | 0.24                            | 0.26                                     | 0.11                             | 0.53*      | 0.44†      | 0.23               |
| Lateral     | 0.67*                           | 0.69*                                    | 0.65*                            | 0.55*      | 0.69*      | 0.66*              |
| Overall     | 0.78*                           | 0.84*                                    | 0.74*                            | 0.63*      | 0.65**     | 0.62*              |

**Diagnostic performance of US in recognising inflammatory arthritis compared with gold standard histology**

For inflammatory arthritis recognition, US B mode had a sensitivity of 42% (95% confidence interval (CI) (13.77%; 69.56%)), a specificity of 90% (95% CI (71.41%; 108.6%)), a negative predictive value (NPV) of 56.25% (95% CI (31.94%; 80.56%)) and a predictive positive value (PPV) of 83.3% (9% CI 5 (33.51%; 113.2%)) compared with histological gold standard.

For inflammatory arthritis recognition, US Doppler mode had a sensitivity of 10% (95% CI (8.59%; 28.59%)), a specificity of 91.16% (76.03%; 107.3%), an NPV 57.14%.
(35.98%; 78.31%) compared with standard vascularity histological assessment.

**DISCUSSION**

Our study is the first to assess correlations for knee synovitis, between US evaluation and histological findings for both B mode and Doppler examination. To date, no study has demonstrated the correlation between B mode semiquantitative synovitis score and both histological H&E stained slides and CD68 staining findings. Given the existing variability of US and histological findings between compartments within the same joint, correlations were not always significant at the compartment level. However, when combined in median scores, moderate to strong correlations were found.

In the literature, controversial results have been described. Anandarajah *et al.* found no significant correlation between US findings and histological findings for synovitis in 15 patients with RA. However, the patients were in remission and the synovial tissue was retrieved during arthroplasty, which may reflect a different, more long-standing disease phenotype. Such synovial tissue may differ from that retrieved in patients with early arthritis, and in our cohort, 70% of the patients (18/26) were untreated at the time of arthroscopy, and the duration of the symptoms ranged from 6 weeks to 6 months.

### Table 4  Correlation for ultrasonography, arthroscopic, histological and immunohistological findings between the three major compartments of the knee

| Ultrasonography | Synovitis grade (r) | Thickness (r) | Power Doppler grade (r) |
|-----------------|---------------------|---------------|-------------------------|
| Superior and lateral | 0.62**             | 0.65***       | 0.15                    |
| Superior and medial   | 0.61*              | 0.48*         | 0.15                    |
| Medial and lateral    | 0.75***            | 0.73***       | 0.44*                   |
| Superior and overall  | 0.75***            | 0.87***       | 0.47*                   |
| Medial and overall    | 0.86***            | 0.87***       | 0.71***                 |
| Lateral and overall   | 0.82***            | 0.80***       | 0.64**                  |

| Arthroscopy | Synovitis (r) | Vascularity (r) |
|-------------|---------------|-----------------|
| Superior and lateral | 0.60*         | 0.51*           |
| Superior and medial   | 0.31         | 0.24            |
| Medial and lateral    | 0.44*         | 0.58*           |
| Superior and overall  | 0.79***       | 0.62*           |
| Medial and overall    | 0.85***       | 0.83***         |
| Lateral and overall   | 0.69***       | 0.75***         |

| Histology | Inflammation (r) | Lining layer thickness (r) | Vascularisation (r) |
|-----------|------------------|---------------------------|---------------------|
| Superior and lateral | 0.47†          | 0.78*                      | 0.64*               |
| Superior and medial   | 0.69*          | 0.69*                      | 0.44†               |
| Medial and lateral    | 0.68*          | 0.95***                    | 0.51†               |
| Superior and overall  | 0.82***        | 0.89***                    | 0.89***             |
| Medial and overall    | 0.93**         | 0.94***                    | 0.74***             |
| Lateral and overall   | 0.70*          | 0.90***                    | 0.79***             |

| Immunohistochemistry | CD68 (r) | FVIII (r) |
|----------------------|----------|----------|
| Superior and lateral | 0.28     | 0.83*    |
| Superior and medial  | 0.49     | 0.71†    |
| Medial and lateral   | 0.34     | 0.61     |
| Superior and overall | 0.85**   | 0.85**   |
| Medial and overall   | 0.82***  | 0.83**   |
| Lateral and overall  | 0.78**   | 0.73*    |

†P<0.1; *P<0.05; **P<0.001; ***P<0.0001.

FVIII, Factor VIII staining; r, correlation coefficient.
Schmidt et al, described association between US PDS and presence or absence of histological pannus. Walther et al, showed a strong correlation (r=0.89; P<0.01) between power Doppler semiquantitative score and the histological qualitative score given by the pathologist after FVIII staining. However, in another study involving 44 subjects using power Doppler evaluation on several joints followed by biopsy, no correlation was found between global inflammation score or blood vessel proliferation. Andersen et al showed a correlation between Doppler colour fraction and overall synovitis score, CD68, CD3, von Willebrand factor in wrists and finger joints, despite finding.

Kelly et al showed a good correlation between PDS, histological vascular area and angiogenic factors. In our work, in respect of vascularity, the correlation between power Doppler score and FVIII staining was not statistically significant. This may be due to a lack of power due to the relatively small cohort. Moreover, only a few patients (4/26, 15.4%) had a positive Doppler activity. This might be explained by the known limited sensitivity of Doppler mode for relatively deep anatomic areas, such as knee joints.

A strong correlation was found between power Doppler grade of the synovitis and histological inflammation score, lining layer hyperplasia score and vascularity score. Our study assessed the correlation between US, arthroscopic and histological findings in the three major compartments of the knee joint. This is the first study assessing the correlation between histological features of the synovial membrane in three well-defined different areas of the knee joint. Indeed, all parameters in each compartment had a moderate to strong correlation with median overall scores. This was also true between LC, MC and SC for US B mode synovitis and measured synovial thickness, for histology and for arthroscopy (although not significant for the SC and MC). However, correlations between MC, LC and SC for CD68 or FVIII staining were not statistically significant.

Our findings suggest that in comparative studies of imaging methods using histology as the reference standard synovial biopsies need to be retrieved from the three major compartments of the knee as the correlations strengthen when each compartment score is compared with the overall median score. In individual compartments, the lack of correlation in some findings could be related to the variability of the findings at the compartment level.

Our work used two different and complementary approaches. After assessing correlations, we explored diagnostic performances of US to detect inflammatory arthropathies using histology as the gold standard. Interestingly, the specificity of US was approximately 90% for both B mode and Doppler mode compared with histological assessment of both inflammation and vascularity. This re-ensures the use of US in detecting synovitis for both diagnosis and response to treatment assessment. However, 10% of the patients having a normal US examination in our cohort still exhibit histological signs of inflammation in their synovial biopsies.

We acknowledge that our work has limitations, the small number of patients in our prospective cohort is one of them. To address this issue, we used non-parametric correlation tests after confirming with professional statistician that results obtained will be generalisable. Indeed, the severity of synovitis in the analysed sample was in general rather low.

In conclusion, we have demonstrated that US examination with both B mode and power Doppler reflects accurately histological inflammation and vascularisation, as well as commonly used CD68 staining in knee synovitis.

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