Further evidence that a cartilage-pannus junction synovitis predilection is not a specific feature of rheumatoid arthritis

L A Rhodes, P G Conaghan, A Radjenovic, A J Grainger, P Emery, D McGonagle

Background: Qualitative differences in synovitis between the cartilage-pannus junction (CPJ) region and the adjoining suprapatellar pouch (SPP) have been reported in rheumatoid arthritis (RA) and spondyloarthropathies. Qualitative differences in synovitis between the cartilage-pannus junction (CPJ) region and the adjoining suprapatellar pouch (SPP) have been reported in rheumatoid arthritis (RA), possibly indicating that the CPJ has a fundamental role in the pathogenesis of RA. Our earlier work in RA and spondyloarthropathies (SpA) has shown synovitis is not uniform within the SPP but greater at the CPJ, suggesting common immunological mechanisms related to a cartilage directed immune response or common biomechanical factors at the pole of the patella determining the greater severity of synovitis.

Osteoarthritis (OA) refers to a heterogeneous group of disorders with common clinical and pathological features; it is strongly associated with aging, is the most prevalent joint disease, and has a strong biomechanical basis. Although synovial inflammation in OA appears to be a secondary process, synovitis is still common; however, there are limited publications describing magnetic resonance imaging (MRI) measurements of synovitis in OA. The purpose of this study was to determine whether the distribution of synovitis between the CPJ region and a distant region in the SPP was the same in OA as in the inflammatory arthropathies. Demonstration of a greater degree of synovitis at the CPJ in OA would support the concept that CPJ synovitis is not unique in RA, but due to common biomechanical and immunological factors in operation, even in degenerative disease.

METHODS

Twenty subjects with established OA of the knee were recruited consecutively. All subjects fulfilled the American College of Rheumatology criteria (clinical and radiographic) for OA. The study protocol was approved by the local ethical review committee, and all patients gave informed written consent to their participation in the study.

Magnetic resonance imaging and image processing

MRI of the knee was performed using a Philips 1.5 T Gyroscan ACS-NT whole body scanner (Philips Medical Systems, Best, the Netherlands) with a Philips quadrature knee coil; the knee was placed in the supine position. The dynamic, contrast enhanced MRI (DEMRI) sequence acquired 40 spoiled T1 weighted gradient echo images from each of five sagittal slices before, during, and after the bolus injection of the contrast agent gadolinium diethylenetriamine pentaaeetic acid (Gd-DTPA). The images were obtained using a repetition time of 11 ms, an echo time of 5.3 ms, and a flip angle of 60° that allowed the acquisition of 5 mm thick slices and a 5 mm gap between slices. The temporal resolution of the DEMRI measurements was 6 seconds for the individual images that constitute the DEMRI dataset. Gd-DTPA was administered at a dose of 0.1 ml/kg body weight 1 second after the acquisition of the first image in the dynamic series.

Commercial software (Analyze – Mayo Clinics, New York) and software developed in-house were used to calculate values of maximal enhancement (ME) and the initial rate of contrast enhancement (IRE) on a pixel by pixel basis; these were displayed as colour overlays on conventional images showing anatomy (fig 1).

Measurements of MR parameters within the synovium at the CPJ and proximal SPP

Measurements were made on the midline sagittal slice at two sites as previously described. Two rectangular regions of interest were defined: one at the superior pole of the patella positioned alongside the CPJ and the second at a remote site in the proximal SPP. Each region had identical superior-inferior dimensions (15 pixels) but variable anterior-posterior dimensions to accommodate the variable synovitis thickness in the sagittal plane (fig 1).

Osteophyte scoring

Unlike early OA and early SpA, where no CPJ region osteophytes were noted, not surprisingly osteophytes were noted in chronic OA; the effect of these on CPJ region synovitis was determined (fig 2). An experienced radiologist.

Abbreviations: CPJ, cartilage-pannus junction; DEMRI, dynamic, contrast enhanced MRI; Gd-DTPA, gadolinium diethylenetriamine pentaaeetic acid; IRE, initial rate of contrast enhancement; ME, maximal enhancement; MRI, magnetic resonance imaging; OA, osteoarthritis; RA, rheumatoid arthritis; SpA, spondyloarthropathies; SPO, superior pole patella osteophyte; SPP, suprapatellar pouch
Statistical analysis
A paired *t* test was used to compare differences between the CPJ and the SPP, and independent *t* tests for differences between the CPJ and the SPO-CPJ. The intraobserver reproducibility for the delineation of regions of interest was calculated using intraclass correlation coefficients and found to be greater than 0.9, as had been observed in the previous study.1

RESULTS
DEMRI parameters at the CPJ and SPP
A significantly larger area of synovitis was evident adjacent to the CPJ than the SPP, representing up to a sixfold difference in some cases. Both the IRE and the ME measures were significantly greater at the CPJ than at the SPP. Overall, these measures correlated with the observation that the synovial area at the CPJ was greater than at the SPP (table 1).

Effect of superior pole patella osteophytes
Radiological scoring highlighted SPOs ranging in size in several patients. Owing to the small number of patients, independent *t* tests did not show statistical significance; mean area values showed less synovitis at the CPJ in patients with SPOs, values of IRE and ME followed the same trend.

### Table 1

<table>
<thead>
<tr>
<th></th>
<th>CPJ v SPP</th>
<th>CPJ v SPO-CPJ</th>
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<tbody>
<tr>
<td><strong>Area</strong> (mm²)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>CPJ</td>
<td>170.00 (p = 0.0001)</td>
<td>238 (p = 0.056)</td>
</tr>
<tr>
<td>SPP</td>
<td>80.00 (p = 0.108)</td>
<td>132 (p = 0.252)</td>
</tr>
<tr>
<td><strong>ME</strong></td>
<td></td>
<td></td>
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<tr>
<td>CPJ</td>
<td>353.25 (p = 0.0001)</td>
<td>473.56 (p = 0.108)</td>
</tr>
<tr>
<td>SPP</td>
<td>148.21 (p = 0.056)</td>
<td>293.221 (p = 0.252)</td>
</tr>
<tr>
<td>SPO-CPJ</td>
<td>293.221 (p = 0.108)</td>
<td>293.221 (p = 0.252)</td>
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<tr>
<td><strong>IRE</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CPJ</td>
<td>3.38 (p = 0.0004)</td>
<td>4.87 (p = 0.252)</td>
</tr>
<tr>
<td>SPP</td>
<td>1.26 (p = 0.108)</td>
<td>2.84 (p = 0.252)</td>
</tr>
<tr>
<td>SPO-CPJ</td>
<td>1.26 (p = 0.108)</td>
<td>2.84 (p = 0.252)</td>
</tr>
</tbody>
</table>

*The initial rate of enhancement (IRE) and maximal enhancement (ME) are shown in arbitrary units.
DISCUSSION

The purpose of this study was to determine whether the distribution of synovitis between the CPJ region and a distant region in the SPP was the same in OA as in the inflammatory arthropathies, where we have already reported a predilection for synovitis at the CPJ in both RA and the SpA.1

The results presented here suggest that the magnitude of synovitis at the CPJ is not disease-specific and applies across the spectrum of degenerative disease as well as inflammatory disease. In addition to investigating synovial areas, this study also assessed other MRI measures of synovitis in the CPJ and SPP. Rates of contrast enhancement in RA have been shown to correlate with both inflammatory activity and blood vessel numerical density.10,11 We noted that IRE and ME measures, MRI surrogates of vascularity, were greater at the CPJ region. The consistent findings using different MRI measures all suggest that the phenotypic expression of synovitis seems to be dictated by common joint factors (biomechanical and immunological) across the spectrum of the rheumatic diseases rather than by specific CPJ features in RA.

MRI is complementary to arthroscopy for the assessment of synovitis. Although arthroscopy allows qualitative and biological assessment of synovitis by histology, it provides little information about the depth or volume of functional characteristics of synovitis. In this regard the ability of MRI to estimate the overall degree and distribution of synovitis may provide additional useful information. Although, MRI measures of synovitis correlate with histological grades, this was not formally assessed in the present study; but the findings of greater synovitis in OA in addition to that already reported in inflammatory arthritis at the CPJ, support the concept that synovitis at that site is not specific for RA, but represents a common joint response across all arthritides.

Although the differences between patients with osteoarthritis and those without must be viewed with some caution owing to the small numbers, the results show that their presence reduces the quantity of synovitis at the CPJ. The importance of this observation is not clear, but it may be that osteoarthritis affects the progression of synovitis at the CPJ by displacing synovial tissue from that site, thus affecting the distribution within the SPP.

In conclusion, this work reaffirms that synovitis is not a feature of OA rather than by specific CPJ features in RA.

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