Psoriatic arthritis (PsA) has a distinctive set of radiographic features that help differentiate this condition from rheumatoid arthritis (RA) and other inflammatory arthropathies. The literature concerning the magnetic resonance imaging (MRI) features of PsA is sparse. While bone erosions and synovitis seem similar to their equivalents in RA [although the distal interphalangeal (DIP) joints may be involved], other spondyloarthropathic features such as dactylitis, periostitis, and enthesitis have been less well described and provide an additional layer of complexity. With the advent of highly effective biologic therapies, it has recently become possible to influence disease progression in PsA, and there is now a need to quantify changes in articular inflammation and damage so that the efficacy of such new treatments can be accurately assessed. The RA MRI score (RAMRIS) was developed through an OMERACT iterative process to capture disease progression in RA. There are currently no validated scoring systems for MRI in peripheral PsA, so we propose to develop one using the same procedure, acknowledging that there will be constraints imposed by the imaging modality itself and the data sets available.

In this initial exercise, we scored images from a preexisting PsA MRI dataset, using a system based on a RAMRIS framework but with additional categories to include PsA-specific features such as extracapsular inflammation. The aim of our project was to determine the pathological features with the greatest interreader reliability for inclusion in a preliminary PsA MRI score (PAMRIS).

**MATERIALS AND METHODS**

The OMERACT MRI group began development of PAMRIS after consensus meetings at American College of Rheumatology 2004 and European League Against Rheumatism 2005. Synovitis and bone erosions were scored as in the RAMRIS system (0–3 and 0–10, respectively). Bone edema was scored 0–3 and in addition, categorized as subchondral, enthesal, or diaphyseal.

Extracapsular inflammation was scored as absent or present (0 or 1). Tendinopathy was also evaluated: tenosynovitis (0–3), intratendinous edema/enhancement (0–1), and edema/enhancement at insertion (0–1).

While it was recognized that other features such as periostitis, bony proliferation, and ankylosis might also be present on some scans, no attempt was made to score these in this first exercise as they were very infrequent.

An image set of MRI scans from 10 PsA patients was chosen (by Charlotte Wiel, Copenhagen). These included images of the 2nd-5th fingers [MCP, proximal interphalangeal (PIP), and DIP joints] obtained on a 0.6 T Philips Panorama MRI unit using the following sequences: 3-D T1 weighted...
RESULTS

Interreader reliability. When mean scores were compared between the 4 readers, there were no significant differences for synovitis, bone edema, tenosynovitis, or extracapsular inflammation. However, Reader 1 scored higher than the others for bone erosions (p < 0.001; Table 1). When this reader was excluded, there was no significant difference between scores for the other readers (p = 0.32). Data for reliability of erosion scores were therefore analyzed both as 3-reader and 4-reader scores.

Table 2 shows single-measures interreader ICC for all components of the score. While the 3-reader ICC for bone erosion (0.91) was very good, and for bone edema was moderate (0.63), interreader reliability for synovitis, tenosynovitis, and extracapsular inflammation was low.

Difficulties for readers. Several readers recorded difficulty in assessing damage and inflammation at the very small joints (PIP and DIP) where image resolution was sometimes poor. It was also felt that recording patterns of bone edema was too difficult, as there was little to separate “subchondral” from “entheseal” in many cases and “diaphyseal” was very rare. Readers also recorded problems defining the exact location of entheseal regions adjacent to small finger joints making the category “extracapsular inflammation” difficult to assess. This was also felt to overlap with the category “edema/enhancement at tendon insertion.” When synovitis coexisted with entheseal and other extracapsular inflammation, with all areas showing increased signal on STIR and post-Gd T1 weighted images, it was also difficult to allocate separate scores for each at the small joints.

While there were examples of florid flexor tenosynovitis within the image set that were recognized by all readers (Figure 1), overall scoring of tendinopathy was only fair, possibly because of difficulties differentiating periostitis from tendinopathy on sagittal STIR images.

DISCUSSION

This preliminary multireader exercise has indicated that creating an MRI scoring system for PsA is possible, with moderate to very good interreader reliability for bone inflammation and damage features (bone edema and erosion). However, difficulties were encountered in assessing synovitis, especially at the very small PIP and DIP joints, and this was apparent from lower interreader reliability than has been achieved using the RAMRIS system in RA. Reliability was also low for scoring extracapsular inflammation and tendinopathy. These difficulties were expected, as clear definitions for the new PsA pathologies had not been provided to readers and there was no pre-exercise reader calibration to optimize their recognition. The scoring sheet used was deliberately overinclusive to capture a wide range of pathologies, with the intention that it could be refined later to incorporate only those features that could be reliably recognized and scored. This may sometimes mean omitting features that are important pathologically but are not well seen on peripheral MRI, as was the case for cartilage involvement, which was omitted during development of RAMRIS.

Another aspect of designing an MRI scoring system for PsA is deciding on the optimal sequences and acquisitions to

Table 1. Scores* (mean and range) for synovitis, bone edema, bone erosions and extracapsular signs of inflammation from 10 PsA MRI scans of the fingers (MCP, PIP, and DIP joints).

<table>
<thead>
<tr>
<th></th>
<th>Synovitis</th>
<th>Bone Edema</th>
<th>Bone Erosion</th>
<th>Extracapsular Inflammation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reader 1</td>
<td>8.1 (6)</td>
<td>1.0 (1.9)</td>
<td>12.7* (4.5)</td>
<td>1.4 (2.1)</td>
</tr>
<tr>
<td>Reader 2</td>
<td>6.2 (5)</td>
<td>2.4 (2.9)</td>
<td>4.7 (4.6)</td>
<td>0.6 (0.7)</td>
</tr>
<tr>
<td>Reader 3</td>
<td>6.7 (5.4)</td>
<td>0.9 (1.5)</td>
<td>2.5 (3.4)</td>
<td>0.8 (1.1)</td>
</tr>
<tr>
<td>Reader 4</td>
<td>7.1 (5.3)</td>
<td>1.2 (2.6)</td>
<td>2.0 (3.2)</td>
<td>1.2 (1.5)</td>
</tr>
</tbody>
</table>

* No significant differences in scores between readers except for erosions, where Reader 1 scores were higher (p < 0.001).
This is more complex than in RA for a number of reasons. There are a broader range of tissues and sites of potential pathology in PsA, but at the same time often fewer joints involved per patient, and these may be asymmetrically distributed. When deciding which pulse sequences to use, spatial resolution and signal-to-noise ratio issues need consideration. T1 weighted SE sequences with fat suppression, pre- and post-contrast with IV gadolinium, are appropriate for most bony lesions including erosions (and were used in this exercise). However, experience in ankylosing spondylitis has suggested that axial T1 weighted sequences lack sensitivity for detection of syndesmophytes and this might also apply to imaging the proliferative bony lesions of peripheral PsA. STIR and T2 weighted fast spin-echo fat-suppressed sequences are very effective for imaging bone edema and soft tissue inflammation, but T2 weighted images are time-consuming to acquire and therefore susceptible to movement artefacts. In this exercise, sagittal STIR sequences of the fingers were useful in identifying dactylitis, usually due to tenosynovitis, as they captured images of the complete ray from the MCP joint to the fingertip. In some cases, false positives could have been scored for tenosynovitis due to periostitis causing increased signal on the inner aspect of the tendon sheath.

To develop PAMRIS further we now intend to focus on the “poorly recognized” categories identified in this exercise, such as extracapsular inflammation and tendinopathy, and decide on definitions, both in terms of anatomic localization and typical MRI signal characteristics. Agreement on image acquisition, sequences, and planes of imaging needs to be reached and future exercises should include pre-exercise reader calibration. From this beginning we hope to refine the process and eventually develop a scoring system that conforms to the OMERACT principles of truth, discrimination, and feasibility.

REFERENCES