Bone resorbing activity in synovial fluids in destructive osteoarthritis and rheumatoid arthritis

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SUMMARY The synovial fluids of patients with a destructive form of osteoarthritis (DOA) were shown to contain high levels of bone resorbing activity as judged by the ability of the fluid to stimulate the release of 45Ca from labelled cultured mouse calvariae. The activity was lost on extended storage of the synovial fluids and was dependent for its effect on cellular activity in bone. Bone resorbing activity was present in most synovial fluids from patients with DOA and rheumatoid arthritis (RA) but occurred at higher levels in the former. In contrast, interleukin 1 (IL1) activity, measured by the mouse thymocytes costimulation assay, was higher in RA than DOA synovial fluids. Little or no bone resorbing or IL1 activity was detected in synovial fluids from patients with pseudogout or non-destructive osteoarthritis. These results suggest that most DOA synovial fluids contain a bone resorbing factor other than IL1. It is considered that this factor may be produced by synovial cells stimulated by hydroxyapatite crystals.

Key word: interleukin 1.

Osteoarthritis is a common disease characterised by destruction of articular cartilage with increased activity and remodelling of subchondral bone. A few patients have the disease in a more destructive form so that, in addition to extensive cartilage loss, there is radiological evidence of bone destruction. Clinical features in these patients may include pain, large cool effusions, and restriction of joint movement, with rapid progression of the destructive process. The synovial fluids of such patients are sterile and free from inflammatory cells, but they contain large amounts of hydroxyapatite crystals. Various names have been used to describe this destructive form of osteoarthritis (DOA), including apatite associated destructive arthropathy,1 Milwaukee shoulder syndrome,2 3 basic calcium phosphate crystal deposition disease,4 5 and analgesic arthropathy.6 7

The mechanisms of joint destruction in osteoarthritides are unknown, though some potential mediators have been investigated. Halverson and colleagues reported that some synovial fluids from patients with DOA contain high levels of activated collagenase and proteases9 and that these enzymes are released from cultured synoviocytes stimulated by hydroxyapatite crystals.8 Others have shown that when cultured synovium engulfs hydroxyapatite crystals there is release of prostaglandin E2,9 a potent bone resorbing agent. If these reactions occur in the patients' joints then they appear to be independent of cellular inflammation, as assessed by histological examination of the patients' synovial membranes.10

Recently a number of bone and cartilage resorbing agents have been described, at least one of which, IL1,11−13 has been detected in the synovial fluids from patients with inflammatory arthritides, especially RA14 15 and gout.16 17 IL1 is known to stimulate synovial cells to release destructive enzymes in vitro.18 19 These findings prompted us to examine synovial fluids from patients with DOA for bone resorbing and IL1 activity.

Patients and methods

Patients Details of the patients are shown in Table 1. Patients studied were those attending rheumatology outpatient clinics at the Bristol Royal Infirmary and requiring aspiration of synovial fluid from a knee or shoulder for diagnostic or therapeutic purposes. A diagnosis of rheumatoid arthritis was made according to the classical American
Rheumatism Association criteria, or of osteoarthritis (OA) by the presence of typical clinical and radiological features. Those with OA were further subdivided into three groups as follows on the basis of synovial fluid and radiological findings: (a) Destructive or apatite associated OA. These patients had radiological evidence of bone destruction and abundant particles staining with alizarin red in the synovial fluid. The clinical and radiological features of this group have been described previously. (b) Simple OA. These patients had no radiological evidence of bone destruction or chondrocalcinosis and no crystals in their synovial fluid, but did have painful joints with radiological evidence of joint space narrowing and subchondral bone reaction. (c) Pyrophosphate arthropathy (PA). These patients had radiological evidence of chondrocalcinosis, synovial fluid pyrophosphate crystals, a hypertrophic bone response, and no destruction of bone.

**SYNOVIAL FLUIDS**

All fluids were tested immediately for the presence of inflammatory cells, and of crystals as shown by alizarin red staining and polarised light microscopy. The fluids were then centrifuged and the supernatants assayed for bone resorbing and IL1 activity.

**BONE RESORPTION ASSAY**

Bone resorbing activity was assessed quantitatively using the mouse calvarial system. Briefly, the skeletons of 1–2 day old mice were labelled with $^{45}$Ca by subcutaneous injection of 37 kBq of $^{45}$CaCl$_2$ (Amersham International) in each mouse. Four days later half calvariae were removed and cultured on stainless steel grids in 1.5 ml modified Bigger's medium (Imperial Laboratories), supplemented with 5% heat inactivated rabbit serum (Wellcome). After 24 hours preincubation bones were transferred either to control medium or to medium containing synovial fluid and cultured for a further 48 hours at 37°C. Aliquots (0.5 ml) of medium were removed and their radioactivity measured by scintillation counting. The results were expressed as the ratio of $^{45}$Ca released into the medium from treated bones to that from control bones; a ratio greater than one indicates the presence of resorptive activity in the synovial fluid. For each experiment positive and negative controls were employed. For positive controls, either vitamin A 22 IU/ml $^{23}$ (Sigma) or prostaglandin E$_2$ $10^{-6}$ mol/l $^{24}$ (Sigma) was used to stimulate bone resorption. For negative controls, dead calvariae were used. These were prepared by exposing calvariae to three cycles of freezing and thawing.

Table 1  **Source of synovial fluid specimens**

<table>
<thead>
<tr>
<th>Type of arthritis</th>
<th>DOA</th>
<th>OA</th>
<th>RA</th>
<th>PA*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of patients</td>
<td>25</td>
<td>10</td>
<td>15</td>
<td>11</td>
</tr>
<tr>
<td>Sex (M:F)</td>
<td>5:20</td>
<td>4:6</td>
<td>5:10</td>
<td>7:4</td>
</tr>
<tr>
<td>Age, mean</td>
<td>72 (57-87)</td>
<td>69 (60-78)</td>
<td>54 (32-73)</td>
<td>63 (46-80)</td>
</tr>
</tbody>
</table>

*PA*= pyrophosphate arthropathy.

**Fig. 1**  **Effect of 5% SF from patients with DOA, 22 IU/ml vitamin A (Vit A), 2 IU/ml IL1, and $10^{-6}$ M PGE$_2$ on calcium release by live mouse calvariae compared with the effect of medium alone (CON). The effect of SF on dead calvariae is also shown. Each experiment was done in triplicate; each result represents the mean of six experiments. Limits represent 1SD. The effect is expressed as the amount of $^{45}$Ca released into the medium. Shaded columns represent release of calcium from dead control and SF treated bones.
thawing. At the end of the experiment some treated bones and their controls were fixed in formalin, decalcified, and processed for histological study. Sections were stained with haematoxylin and eosin and examined by light microscopy.

**INTERLEUKIN 1 ASSAY**

This was performed as described elsewhere, with some modifications. Quadruplicate cultures were prepared in microtest well plates (Nunc); each contained 2×10^5 thymocytes from C3H/HeJ mice in 200 μl of RPMI 1640 medium containing 1% phytohaemagglutinin (PHA, Gibco) and dilutions of synovial fluid (SF). The SF samples were treated with sheep testis hyaluronidase 100 U/ml (Sigma) before testing. Control wells contained thymocytes alone or thymocytes and PHA. As a positive control, thymocytes were cultured in medium containing 0.25 U/ml purified IL1 (Genzyme). After incubation for 48 hours at 37°C the thymocytes were pulsed with 37 kBq/well of [3H]thymidine (Amersham) for the last six hours and harvested with an automatic cell harvester. Thymidine uptake was measured in a liquid scintillation counter. Data were expressed as the mean (SD) of quadruplicate cultures.

**STATISTICAL ANALYSIS**

Student's t test was used to compare the control and experimental groups in both assays.

**Results**

**BONE RESORBING ACTIVITY OF DOA SYNOVIAL FLUID**

Fig. 1 shows that labelled calvariae cultured with DOA synovial fluid released more radioactivity than calvariae cultured in medium alone (p<0.001). The known bone resorbing agents, vitamin A, IL1, and
PGE₂, induced a similar response. In contrast, SF failed to stimulate the release of calcium from dead bones.

It was noticed that SF appeared to have reduced bone resorbing activity on storage. To verify this observation samples were stored at −70°C and repeatedly thawed for each testing at intervals. Fig. 2 shows the result of experiment with three fluids. In each case there was a progressive loss of activity with time. In a further experiment (not illustrated) aliquots of the initial sample were obtained at the time of initial storage and only one sample thawed at each time. Activity again diminished with time, which suggests that the loss of activity is associated with length of time of storage rather than with the effect of repeated thawing and refreezing. Because of this all subsequent experiments on bone resorbing activity were conducted on fresh synovial fluids.

Experiments were set up to test the effect of varying the concentration of synovial fluid on bone resorbing activity. Synovial fluids with bone resorbing activity from patients with DOA were diluted and added to live labelled bones. Fig. 3 shows that there was stimulation of bone resorption at the lowest concentration tested, which was 1%. Peak resorption occurred at concentrations between 5% and 7%. At levels above 10% no further enhancement of calcium release was found; on the contrary, high concentrations inhibited bone resorption.

To test if the reduced effect at high concentrations of synovial fluid was due to a toxic effect on the cultured bone, high concentrations of SF were added to cultures stimulated with vitamin A. Whereas the ratio of ⁴⁵Ca released from vitamin A treated bones to that released from control bones was 1-6, the addition of 20% SF from patients with DOA reduced the ratio to less than 1 (complete inhibition). Suboptimal concentrations of SF (1-2%) had no effect on vitamin A induced ⁴⁵Ca release.

Fig. 4 shows a comparison between the levels of bone resorbing activity in DOA synovial fluids and those in SF from other arthritides. High levels of activity were present in 24/25 of DOA synovial fluids, whereas little or no activity was detected in fluids from patients with OA and PA. Most (12/15) RA fluids were active, but the average level was significantly less than that found in DOA (p<0-05). This could not be attributed to variations in the assay as the amount of ⁴⁵Ca release induced by vitamin A, the positive control in these experiments, fell within the range shown for vitamin A in Fig. 1. The possibility that this difference resulted from the RA fluids being tested at a suboptimal concentration was investigated by employing different concentrations of RA fluids. The results showed that, as with DOA, maximal activity occurred at a concentration of 6%. The three RA synovial fluids which were inactive in the initial test (Fig. 4) showed no activity at any of the dilutions tested.

**Histo logical examination**
The results recorded in Fig. 1 show that calcium release from bone induced by SF depends on live cells in the bone, presumably osteoclasts. To investigate this, histological sections were prepared from living bones which had been incubated with either synovial fluid or with medium alone. Although there appeared to be more osteoclasts in
the former group (Fig. 5b) than in the latter (Fig. 5a), multinucleated osteoclasts were seen in the latter group. Thus such histological observations should be treated with some caution.

**INTERLEUKIN 1 ACTIVITY OF DOA AND OTHER SYNOVIAL FLUIDS**

The effect of different concentrations of DOA and RA synovial fluids on thymocyte proliferation was measured. Fig. 6 shows that fluids from both groups of patients gave maximum proliferation at a concentration of 3%. Although the curves for both groups were similar, the fluids from patients with DOA induced less proliferation than those from patients with RA.

Fig. 7 compares the thymocytes co-mitogenic stimulating activity of fluids from patients with DOA, OA, RA, and PA, all tested at a concentration of 3%. The results show that synovial fluids from patients with OA and PA lacked activity, whereas fluids from most patients with RA and some with DOA stimulated thymocyte proliferation.

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**Fig. 5** Photomicrographs of calvariae. (Haematoxylin and eosin.) (a) In the control tissue there is solid bone with surrounding mononucleated cells with few if any osteoclasts. (b) In a typical area from tissue treated with SF the bone shows multiple lacunae associated with numerous multinucleated giant cells (arrow).
tion. As a group rheumatoid fluids were significantly more active than those from patients with DOA (p<0.001).

**Discussion**

The main conclusion from this study is that synovial fluid from patients with the rapidly destructive form of osteoarthritis contains one or more bone resorbing agents. The activity of the fluids was comparable in magnitude with that of other known bone resorbing agents such as vitamin A, IL1, and PGE2. Moreover, their mechanism of action appears similar in some respects. As the fluids failed to cause release of calcium from dead bones their mode of action cannot be due to direct dissolution of hydroxyapatite crystals but must be due to the stimulation of cells in the bone, which then modified...
resorption. A similar dependence on the presence of healthy live osteoclasts (and osteoblasts) has been reported for bone resorption induced by parathormone, osteclast activating factor, and IL1. Further evidence of osteoclast activation was sought in this study by examining the histological appearances of bone incubated with SF and comparing these with the appearances of bone which had been incubated with medium alone. Although increased numbers of large multinucleated osteoclasts seemed to be present in the former group, similar, though fewer, cells were present in the control tissues. Confidence in these findings would require a full morphometric analysis of large numbers of sections.

The bone resorbing activity of SF was most marked at concentrations between 5% and 7%. At higher concentrations the effect was less marked. This could be due to high concentrations of the bone resorbing agent(s) down regulating its own activity, to a specific inhibitor of the agent coming in to play as its concentration increases; or to high concentrations of synovial fluid providing an unfavourable environment for cellular activity in the bone. The fact that high concentrations of DOA synovial fluids prevented vitamin A from exerting its bone resorbing effect provides evidence in support of the last hypothesis.

It was considered that the bone resorbing activity in synovial fluids from patients with DOA might be due to IL1. When these fluids were assayed for IL1 activity using a standard mouse thymocyte mitogenic assay, IL1-like activity was indeed found in some of the fluids, but other fluids which stimulated bone resorption failed to show IL1-like activity. In addition, when fluids from patients with DOA and RA were compared there was a discrepancy between the two types of assay. Fluids from the patients with DOA stimulated more bone resorption than those from patients with RA, whereas the reverse was true for IL1-like activity. Our interpretation of these results is that although IL1 may contribute to the bone resorbing activity in both DOA and RA fluids, a different factor, possibly PGE2, is responsible for most of the bone resorbing activity in DOA. This possibility gains credence from the finding that much of the activity in DOA synovial fluids was dialysable. Indeed, even in RA synovial fluid other bone resorbing factors are likely to be present as recent work has shown that tumour necrosis factor has bone resorbing activity and high levels of tumour necrosis factor messenger RNA have been extracted from RA synovial cells (Buchan G, personal communication).

The destructive form of osteoarthritis is characterised by the presence of hydroxyapatite crystals in the synovial fluid and degeneration of the articular cartilage with exposure of the underlying bone and extensive destruction and resorption of the latter. We have preliminary evidence that hydroxyapatite crystals can stimulate macrophages to release a factor which shows bone resorbing activity but not IL1-like activity. At present it is unclear to what extent the production of such bone resorbing factors can explain the relation between hydroxyapatite crystals and bone resorption in DOA. Current work is directed towards characterising the bone resorbing factor in order to compare it with other known resorbing agents.

This work was supported in part by grants from the Arthritis and Rheumatism Council and MRC. Dr Alwan was in receipt of an overseas research student award. We would like to thank Dr Papoian for his help and advice.

References

Bone resorbing activity in OA and RA


