Immunolocalisation studies on six matrix metalloproteinases and their inhibitors, TIMP-1 and TIMP-2, in synovia from patients with osteo- and rheumatoid arthritis

R M Hembry, M R Bagga, J J Reynolds, D L Hamblen

Progressive degradation of the extracellular matrix of connective tissues and destruction of articular cartilage, intra-articular ligaments, tendons, and subchondral bone are major features of the arthritides, leading to disfigurement and permanent loss of function. Although proteinases of all mechanistic classes have the potential to degrade individual connective tissue macromolecules in vitro, the current view is that the initial step in matrix degradation is often an extracellular proteolytic process involving matrix metalloproteinases (MMPs). MMPs are a family of metal-dependent proteinases, which are active at neutral pH and secreted by mesenchymal and haemopoietic cells as proenzyme forms requiring extracellular activation. Their activities are further regulated by secreted inhibitors, the tissue inhibitors of metalloproteinases (TIMPs). The MMP family can be divided into subgroups, three of which contain multiple gene products. The collagenases specifically cleave interstitial collagens types I, II, and III at a single locus; gelatinases (type IV collagenases) cleave denatured collagens and type IV collagen; while the stromelysins are more general proteinases, cleaving proteoglycan core protein, fibronectin and type IV collagen. In addition, a truncated MMP, matrilysin has a substrate specificity similar to that of the stromelysins. Together, these enzymes have the ability to degrade all the macromolecular components of connective tissues.

There is considerable evidence that connective tissues removed from patients with rheumatoid arthritis secrete greater concentrations of MMPs and their inhibitors into culture media than normal tissues. Cultures of adherent cells derived from rheumatoid synovia secreted collagenses into the culture media and collagenses was detected within such cells and at sites of cartilage erosion in rheumatoid joints by immunolocalisation. Stromelysin 1 and collagenase have been demonstrated by immunolocalisation in lining cells of synovia and also mRNA for these proteins was detected by in situ hybridisation. The last studies also identified collagenase and stromelysin mRNA in osteoarthritic synovium in smaller concentrations than in rheumatoid synovium. Primary rheumatoid synovial fibroblasts expressed high levels of stromelysin 1 mRNA but only small amounts of stromelysin 2 mRNA. MMPs have also been measured in rheumatoid synovial fluids and synovial fluids from osteoarthritic knee joints. TIMP-1 has been detected in lining cells of rheumatoid synovia by immunolocalisation and in situ hybridisation, and measured in rheumatoid synovial fluids and synovial fluids from osteoarthritic knee joints.
Although the presence of collagenase, stromelysin and TIMP-1 have been well documented, other MMPs may also contribute to a predominance, either overall or focal, of proteases over inhibitors, which could result in the breakdown of joint tissues in the arthritides. In this paper we have analysed synovia from seven joints with inflammatory arthritis and three osteoarthritic joints by indirect immunofluorescence microscopy to document the frequencies and distributions of six MMPs (collagenase, stromelysin 1 and 2, gelatinases A and B, matrilysin) and TIMP-1 and TIMP-2, in order to assess the likely importance of each in the arthritic process.

Materials and methods

Tissues

Synovial samples were removed from 10 patients (table 1). Seven of these patients had a diagnosis of rheumatoid arthritis, seropositive rheumatoid arthritis was present in five, and the other two were seronegative but otherwise satisfied the ARA criteria for rheumatoid arthritis. Three of the five patients with seropositive disease (patients 2, 3 and 4) were in an early proliferative phase of their disease showing only minimal damage to the articular cartilage, and were therefore managed by surgical synovectomy. The remaining two patients (1 and 5) had disease at a more advanced stage, with major destruction of both bone and articular cartilage leading to established secondary osteoarthritic changes. In these the surgical management was by joint debridement or arthroplasty and there was only minimal macroscopic evidence of increased vascularity and synovial hypertrophy. The two patients with atypical seronegative rheumatoid arthritis (patients 6 and 7) both showed advanced joint destruction, although this was still associated with marked proliferative synovial activity. The duration of the inflammatory arthritis varied from two to 24 years, but all patients had received prolonged drug treatment with non-steroidal anti-inflammatory agents and some second-line therapy, such as sulphasalazine or gold salts. Only one patient (patient 6) with atypical seronegative arthritis had been taking systemic steroids for several months before removal of the synovial specimen. All joints in patients with rheumatoid arthritis had received one or more local steroid injections, with the exception of the elbow in patient 1 and the wrist in patient 7.

In the three patients with osteoarthritis, synovial specimens were removed at the time of surgery for replacement arthroplasty of the joint. In one of these (patient 8) there was macroscopic and microscopic evidence of a pyrophosphate arthropathy (pseudogout), with areas of stippled calcification in all the articular tissues in addition to marked synovial proliferation.

In the laboratory, small pieces were cut from each synovial sample and cultured for 6 h in Dulbecco’s Modified Eagle’s Medium with 10% fetal bovine serum and 5 μmol/l monensin (Sigma) in an atmosphere of 5% carbon dioxide in air. They were then frozen in liquid nitrogen. A piece adjacent to each cultured sample was also taken and was frozen ex vivo.

Antibodies

Specific polyclonal antibodies to human gelatinase A, pig gelatinase B (which cross reacts with human gelatinase B but not human gelatinase A), human stromelysin (which reacts strongly with stromelysin 1 and weakly with stromelysin 2), human TIMP-1 and TIMP-2 were raised in sheep. The characterisation of these antisera including species specificity, inhibition curves and immunoblotting experiments with purified antigen are detailed in the references cited. Human interstitial collagenase was purified according to the method of Whitham et al., injected into a sheep and the resulting antisera characterised by Western blotting, and by immunolocalisation on NSO mouse myeloma cells transfected with matrilysin. It was shown to be negative on immunolocalisation of NSO mouse myeloma cells transfected with human stromelysin 1, human stromelysin 2, human collagenase or human gelatinase A. Antiser to human stromelysin 2 were raised in rabbits and sheep by injecting either 1 mg or 2 mg, respectively, of the peptide VPDKVSIVPSGEPKAC (residues 257-272 in the proenzyme) conjugated to tuberculin purified protein derivative and emulsified in incomplete Freund’s adjuvant. Booster injections of 0.5 or 1.0 mg, respectively, were given at three-weekly intervals and blood removed 10 days after each injection. Immunoglobulins (IgGs) were prepared from the bleeds with the highest titres. Specificities were determined by Western blotting and immunolocalisation on NSO mouse myeloma cells transfected with stromelysin 2. The antisera did not cross-react.

<table>
<thead>
<tr>
<th>Patient</th>
<th>Sex</th>
<th>Age (yr)</th>
<th>Diagnosis</th>
<th>Joint</th>
<th>Disease duration (yr)</th>
<th>Operation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>F</td>
<td>62</td>
<td>RA (secondary OA)</td>
<td>Elbow</td>
<td>7</td>
<td>Synovectomy and joint debridement</td>
</tr>
<tr>
<td>2</td>
<td>M</td>
<td>69</td>
<td>RA (active)</td>
<td>Knee</td>
<td>3</td>
<td>Synovectomy</td>
</tr>
<tr>
<td>3</td>
<td>F</td>
<td>30</td>
<td>RA (active)</td>
<td>Knee</td>
<td>2</td>
<td>Synovectomy</td>
</tr>
<tr>
<td>4</td>
<td>F</td>
<td>26</td>
<td>RA (active)</td>
<td>Knee</td>
<td>11</td>
<td>Synovectomy</td>
</tr>
<tr>
<td>5</td>
<td>F</td>
<td>51</td>
<td>RA (secondary OA)</td>
<td>Knee</td>
<td>13</td>
<td>Arthroplasty</td>
</tr>
<tr>
<td>6</td>
<td>F</td>
<td>87</td>
<td>Seronegative rheumatoid arthritis</td>
<td>Shoulder</td>
<td>2</td>
<td>Synovectomy and joint debridement</td>
</tr>
<tr>
<td>7</td>
<td>F</td>
<td>65</td>
<td>Seronegative rheumatoid arthritis</td>
<td>Wrist</td>
<td>24</td>
<td>Silastic arthroplasty</td>
</tr>
<tr>
<td>8</td>
<td>F</td>
<td>68</td>
<td>OA (pyrophosphate arthropathy)</td>
<td>Knee</td>
<td>4</td>
<td>Arthroplasty</td>
</tr>
<tr>
<td>9</td>
<td>F</td>
<td>67</td>
<td>OA</td>
<td>Knee</td>
<td>20</td>
<td>Arthroplasty</td>
</tr>
<tr>
<td>10</td>
<td>M</td>
<td>69</td>
<td>OA</td>
<td>Hip</td>
<td>3</td>
<td>Arthroplasty</td>
</tr>
</tbody>
</table>

RA = Rheumatoid arthritis; OA = osteoarthritis.

Table 1: Details of patients studied.
react with stromelysin 1, either on Western blotting with purified antigen or on immunolocalisation of NSO mouse myeloma cells transfected with stromelysin 1.34 Pooled normal sheep serum (NSS) was used as control and IgGs were prepared by ammonium sulphate precipitation.31 The secondary antibody was the fluorescein isothiocyanate conjugated (FITC) pig anti-sheep Fab previously described (pig-FITC).31

IMMUNOLOCALISATION

The MMPs and TIMPs are usually secreted in vivo in small amounts and are not stored intracellularly. To increase sensitivity in order to locate precisely the cellular source of these proteins, tissues were incubated with the ionophore monensin, which blocks translocation of secreted protein while allowing synthesis to continue.31 37 This procedure results in intracellular accumulation of antigen in the Golgi apparatus and secretory vesicles of cells, which can then more readily be localised by indirect immunofluorescence.

Frozen sections (7 μm) were cut on a cryostat and taken onto glass slides. Sections were fixed (4% freshly prepared formaldehyde for 30 minutes), permeabilised (0.1% Triton X-100 for five minutes) and incubated in 4-chloro-1-naphthol (2-8 mMol/l in methanol/phosphate buffered saline (PBS) with 0.01% hydrogen peroxide for 12 minutes) to prevent non-specific binding of fluorescein to eosinophils. Sections were then incubated for 30 minutes at room temperature with IgG preparations of either antisera or NSS (all 50 μg/ml). Twenty four sections were cut at each location in all samples, and the antibodies assigned to each set of eight (for example anticolonagenase on Nos 2, 10, and 18) to limit variation in tissue histology as far as possible. The sections were then washed in PBS three times for five minutes after each of the above steps. They were then incubated for 30 minutes with the pig-FITC second antibody, washed, and counterstained with methyl green 1 mg/ml for two minutes to stain nuclei red.

All sections were mounted in Citifluor (University of Kent at Canterbury) and viewed by fluorescence microscopy on a Zeiss Photomicroscope III with epifluorescence and wide band FITC filter. Photographs were taken on Agfachrome 1000 RS film uprated during processing to 2000 ASA. Some sections were also analysed by confocal microscopy using a Biorad MRC 600 confocal microscope with a krypton/argon mixed gas laser; photographs were taken on Agfa APX 25 film from a colour monitor. Coverslips were then removed and the sections stained with Harris’s haematoxylin and eosin, observed by bright field optics and photographed on Ektachrome 50 film.

Results

Synovia from seven patients with a history of rheumatoid arthritis (patients 1–7, table 1) and three patients with a history of osteoarthritis (patients 8–10) were examined immunohistochemically in detail using specific antisera to six MMPs and TIMP-1 and TIMP-2. The synovia showed a considerable range in histological appearance and MMP distributions (table 2). Stromelysin 1 was observed in all synovia, bound to extracellular matrix, within cells, or both, indicating stromelysin synthesis. Intracellular staining for stromelysin 1, collagenease, gelatinase A, and TIMP-1 was focal, with marked regional variations, but in general the extent of staining correlated with the degree of synovial inflammation. Interesting observations from individual patients are described in detail below.

The synovial samples from patient 7, with seronegative polyarthritis, were heavily infiltrated with inflammatory cells with a well defined lining layer three to five cells thick overlying fibrovascular tissue. When this tissue was cultured in monensin, sectioned, and exposed to MMPs, extensive regions of the lining cells showed strong intracellular immunofluorescence for stromelysin (fig 1a), with immunofluorescence of the extracellular matrix (fig 1a). However, there were also regions where both lining cells and extra-

---

Table 2  Histology and distribution of MMPs and TIMPs in ex vivo synovia and those cultured for six hours with monensin to accumulate intracellular antigen

<table>
<thead>
<tr>
<th>Patient</th>
<th>Synovial histology</th>
<th>Coll.</th>
<th>SL1</th>
<th>SL2</th>
<th>Mat.</th>
<th>Gel. A</th>
<th>Gel. B</th>
<th>TIMP-1</th>
<th>TIMP-2</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Thin synovial lining, little infiltrate</td>
<td>0</td>
<td>□ L+: S++</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>2</td>
<td>Heavily infiltrated proliferative fronds</td>
<td>0</td>
<td>■ S+++</td>
<td>0</td>
<td>0</td>
<td>□ S+</td>
<td>0</td>
<td>□ S+</td>
<td>0</td>
</tr>
<tr>
<td>3</td>
<td>Heavily infiltrated synovial fronds with cuffed blood vessels</td>
<td>0</td>
<td>■</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>4</td>
<td>Fibrovascular tissue with foci of inflammatory cells</td>
<td>S+</td>
<td>■ S+</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>S+</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>Thin synovial lining with some foci of inflammatory cells</td>
<td>0</td>
<td>□ L+</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>PMNL</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>6</td>
<td>Heavily infiltrated synovial fronds</td>
<td>L++</td>
<td>■ L++</td>
<td>0</td>
<td>L++</td>
<td>S++</td>
<td>0</td>
<td>L++</td>
<td>0</td>
</tr>
<tr>
<td>7</td>
<td>Heavily infiltrated synovial fronds</td>
<td>S++</td>
<td>L++</td>
<td>0</td>
<td>S++</td>
<td>L++</td>
<td>0</td>
<td>S++</td>
<td>0</td>
</tr>
<tr>
<td>8</td>
<td>Thin synovial lining with little infiltrate</td>
<td>0</td>
<td>□ L+</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>9</td>
<td>Thin synovial lining with little infiltrate</td>
<td>S++</td>
<td>S++</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>S++</td>
<td>0</td>
</tr>
<tr>
<td>10</td>
<td>Thin synovial lining, a few inflammatory cell foci</td>
<td>0</td>
<td>□ S+</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Coll. = Collagenease; SL1, SL2 = stromelysins 1 and 2; Mat. = matrixsyn; Gel. A, Gel. B = gelatinases A and B; TIMP-1, TIMP-2 = tissue inhibitors of metalloproteinases 1 and 2; L = lining cells; S = stromal cells; PMNL = polymorphonuclear leucocytes.

Immunofluorescence indicating antigen synthesis: + = 1–10 positive cells per section ...... +++ = >50 positive cells. Connective tissue matrix staining: □ = weak, small area .... □... □ = intense, extensive staining.
cellular matrix were negative. Adjacent sections stained with anti-collagenase also had intracellular immunofluorescence in the same lining cell regions where stromelysin was seen (fig 1b) and fibrovascular tissue below the synovial fronds contained many positive stromal cells, but there was no staining of extracellular matrix. Sections stained with anti-gelatinase A had no immunofluorescence of lining cells or matrix, but large numbers of stromal cells in the underlying connective tissue had weak intracellular immunofluorescence (fig 2a). Sections stained with anti-TIMP-1 showed strong immunofluorescence of lining cells and many underlying stromal cells (fig 1c), indicating more widespread cellular synthesis of TIMP-1 than of either of the enzymes. Sections of this synovium were negative for stromelysin 2, matrilysin, gelatinase B, and TIMP-2, and for the control NSS IgG.

Synovium from patient 6, the other subject with atypical seronegative rheumatoid arthritis, was also heavily infiltrated with inflammatory cells and, when stained with anti-stromelysin, showed strong immunofluorescence of extracellular matrix and cells (fig 2b). As the anti-stromelysin antiserum reacts with both stromelysins 1 and 2 (see Methods), adjacent sections were stained with either the rabbit or the sheep antisera to stromelysin 2, and were found to be negative (fig 2c), indicating that the matrix and cellular immunofluorescence were attributable to stromelysin 1. However, sections of this tissue stained with anti-matrilysin contained discrete areas of lining cells with intracellular fluorescence (fig 2d), indicating matrilysin synthesis. These areas frequently coincided with areas of bright stromelysin fluorescence. The distributions of collagenase, gelatinase A, and TIMP-1 were similar to those seen in patient 7, with TIMP-1

Figure 1 Distribution of stromelysin, collagenase and TIMP-1 in a rheumatoid synovium. Synovium from patient 7 was cultured in monensin, frozen, sectioned, and stained by indirect immunofluorescence with antibodies to (a) stromelysin, (b) collagenase and (c) TIMP-1. Sections were analysed on an MRC 600 confocal microscope using the 488 nm line, and data collected by Kalman averaging over 20 scans at slow scan speed, followed by normalising the image before photography. Bars = 100 μm. a: Anti-stromelysin. Intense intracellular fluorescence is present in synovial lining cells, and in some stromal cells (arrows). The extracellular matrix below the lining is also strongly fluorescent, but there are no positive cells (arrow heads). b: Anti-collagenase. Synovial lining cells have strong intracellular immunofluorescence, with only a few stromal cells weakly positive. There is no matrix fluorescence. c: Anti-TIMP. Synovial lining cells and many underlying stromal cells (arrows) have intracellular fluorescence.
Immunolocalisation studies on six matrix metalloproteinases and their inhibitors, TIMP-1 and TIMP-2

Immunolocalisation studies on six matrix metalloproteinases and their inhibitors, TIMP-1 and TIMP-2.

**Figure 2** Immunolocalisation of gelatinase, stromelysin and matrilysin in rheumatoid synovia. Synovia from patients 7 (a), and 6 (c, d) were cultured in monensin before freezing. Synovium from patient 2 (e, f) was frozen ex vivo. Tissues were sectioned and stained with anti-gelatinase (a, f), anti-stromelysin (b, e), sheep anti-stromelysin 2 (c) or anti-matriplysin (d). Sections were observed by immunofluorescence microscopy. Nuclei are counterstained and appear red. Bars = 20 µm. a: Anti-gelatinase. Some connective tissue cells have weak juxtanuclear green fluorescence indicating gelatinase A synthesis. b: Anti-stromelysin. Extracellular matrix of synovial lining and below is strongly fluorescent, indicating matrix-bound stromelysin, and some cells have vesicular intracellular fluorescence of stromelysin synthesised during the culture period. c: Anti-stromelysin 2. The identical region of the section adjacent to (b) had no green fluorescence, showing that the intracellular and extracellular staining in (b) represents stromelysin 1. d: Anti-matriplysin. Some synovial lining and sublining cells have intracellular immunofluorescence, indicating matrilysin synthesis. e: Anti-stromelysin. Synovium ex vivo has bright stromelysin immunofluorescence of connective tissue matrix. This intense and extensive matrix staining also occurred in patients 2, 3 and 6 and is represented by ■ in table 2. The lining cell layer is negative but contains cells with yellow autofluorescent granules, probably phagocytosed material. f: Anti-gelatinase A. The section adjacent to (e) had a small area of matrix immunofluorescence immediately underlying the synovial lining (arrow), representing gelatinase A bound to connective tissue components.

Synovia from patients 7 (a), and 6 (c, d) were cultured in monensin before freezing. Synovium from patient 2 (e, f) was frozen ex vivo. Tissues were sectioned and stained with anti-gelatinase (a, f), anti-stromelysin (b, e), sheep anti-stromelysin 2 (c) or anti-matriplysin (d). Sections were observed by immunofluorescence microscopy. Nuclei are counterstained and appear red. Bars = 20 µm. a: Anti-gelatinase. Some connective tissue cells have weak juxtanuclear green fluorescence indicating gelatinase A synthesis. b: Anti-stromelysin. Extracellular matrix of synovial lining and below is strongly fluorescent, indicating matrix-bound stromelysin, and some cells have vesicular intracellular fluorescence of stromelysin synthesised during the culture period. c: Anti-stromelysin 2. The identical region of the section adjacent to (b) had no green fluorescence, showing that the intracellular and extracellular staining in (b) represents stromelysin 1. d: Anti-matriplysin. Some synovial lining and sublining cells have intracellular immunofluorescence, indicating matrilysin synthesis. e: Anti-stromelysin. Synovium ex vivo has bright stromelysin immunofluorescence of connective tissue matrix. This intense and extensive matrix staining also occurred in patients 2, 3 and 6 and is represented by ■ in table 2. The lining cell layer is negative but contains cells with yellow autofluorescent granules, probably phagocytosed material. f: Anti-gelatinase A. The section adjacent to (e) had a small area of matrix immunofluorescence immediately underlying the synovial lining (arrow), representing gelatinase A bound to connective tissue components.

Synovia from patients 7 (a), and 6 (c, d) were cultured in monensin before freezing. Synovium from patient 2 (e, f) was frozen ex vivo. Tissues were sectioned and stained with anti-gelatinase (a, f), anti-stromelysin (b, e), sheep anti-stromelysin 2 (c) or anti-matriplysin (d). Sections were observed by immunofluorescence microscopy. Nuclei are counterstained and appear red. Bars = 20 µm. a: Anti-gelatinase. Some connective tissue cells have weak juxtanuclear green fluorescence indicating gelatinase A synthesis. b: Anti-stromelysin. Extracellular matrix of synovial lining and below is strongly fluorescent, indicating matrix-bound stromelysin, and some cells have vesicular intracellular fluorescence of stromelysin synthesised during the culture period. c: Anti-stromelysin 2. The identical region of the section adjacent to (b) had no green fluorescence, showing that the intracellular and extracellular staining in (b) represents stromelysin 1. d: Anti-matriplysin. Some synovial lining and sublining cells have intracellular immunofluorescence, indicating matrilysin synthesis. e: Anti-stromelysin. Synovium ex vivo has bright stromelysin immunofluorescence of connective tissue matrix. This intense and extensive matrix staining also occurred in patients 2, 3 and 6 and is represented by ■ in table 2. The lining cell layer is negative but contains cells with yellow autofluorescent granules, probably phagocytosed material. f: Anti-gelatinase A. The section adjacent to (e) had a small area of matrix immunofluorescence immediately underlying the synovial lining (arrow), representing gelatinase A bound to connective tissue components.

The synovium removed from patient 1, with late stage quiescent rheumatoid arthritis, consisted of fibrovascular stroma without lymphocytic foci bordered by a lining layer one or two cells thick. Sections of tissue cultured in monensin then stained with the antiserum to stromelysin contained a few lining cells and some stromal cells with intracellular immunofluorescence, indicating stromelysin synthesis; a small area of extracellular matrix immediately below the lining layer also had immunofluorescence. Adjacent sections stained with either the rabbit or sheep antisera to stromelysin 2 were negative, indicating that the cells were synthesising stromelysin 1. Sections stained with the other antisera were negative, confirming the histological picture of a relatively inactive joint. Adjacent sections stained with NSS IgG as control were also negative. Ex vivo synovium from patient 5, which also showed more advanced disease with secondary osteoarthritic changes, contained a few polymorphonuclear leucocytes with intracellular gelatinase B immunofluorescence within small inflammatory cell foci. Polymorphonuclear leucocytes were identified in other ex vivo synovia using the methyl green nuclear counterstain and confirmed histologically, but they were negative for gelatinase.
B, presumably having degranulated before tissue excision.

As the synovial samples cultured in monensin from all patients had some stromelysin staining of extracellular matrix, samples frozen ex vivo without culture were also examined. Areas of positive matrix (but no positive cells) were again observed in samples from all patients. The synovium from patient 2 was highly villous with many lymphocytic foci and perivascular inflammation, bordered by a thickened lining layer: when frozen ex vivo, sectioned, and stained with anti-stromelysin IgG, sections had bright immunofluorescence of stromal matrix throughout (fig 2e). Adjacent sections stained with NSS, anti-collagenase, or anti-TIMP IgGs were negative, but sections stained with anti-gelatinase A had positive matrix in one limited area only (fig 2f).

These data indicate that stromelysin 1 and gelatinase A were bound to matrix in vivo, not as a result of the six hour period of culture with monensin. Sections from adjacent tissue cultured in monensin for six hours and stained with either NSS or anti-collagenase IgGs were negative, but sections stained with anti-stromelysin IgG had immunofluorescence both on collagenous matrix and within cells just below the lining layer, indicating stromelysin synthesis (not illustrated). Sections stained with anti-gelatinase A IgG had positive matrix only in one small region and a few positive cells (not illustrated). Anti-TIMP-1 stained sections also had a few positive stromal cells, and localised positive matrix mainly just below the lining layer in regions of cells with intracellular stromelysin (not illustrated).

As the immunofluorescence in the anti-stromelysin stained slides was so extensive and striking and does not occur in normal synovia (data not shown), the presence of stromelysin was cross checked by staining sections with an antiserum raised against active N-terminal stromelysin isolated from rabbit bone culture media; this antiserum cross reacts with human stromelysin. Both the anti-human and the anti-rabbit stromelysin antisera immunolocalise both active high molecular mass and latent stromelysin on collagen fibrils. The sections had both intracellular and matrix fluorescence, confirming the widespread distribution of stromelysin in this synovium. The experiments of Allan et al showed that this matrix bound enzyme could be either latent or active stromelysin, but also showed that TIMP-1 binds to matrix bound active stromelysin and not matrix bound prostromelysin. The immunolocalisation of TIMP-1 on matrix in the same region as stromelysin in this synovial sample suggests that at least a proportion of the stromelysin is in the active form. Sections stained with either the rabbit or sheep antisera to stromelysin 2 were negative, indicating that the matrix and cellular staining were attributable to stromelysin 1.

Synovia from the three osteoarthritic joints (patients 8–10) were found to have a distribution of MMP similar to that in patient 1 with inactive rheumatoid arthritis (table 2). All three had small areas of weak stromelysin 1 immunofluorescence on extracellular matrix in both ex vivo and cultured samples. The synovial lining cells were negative for collagenase in all samples, but one patient (patient 8) had a few lining cells positive for stromelysin.

Discussion

In this paper we have documented the frequencies and distributions of six MMPs and their natural inhibitors, TIMP-1 and TIMP-2, in seven rheumatoid and three osteoarthritic synovia. Stromelysin 1 was present in all synovia, in ex vivo tissue bound to extracellular matrix, within cells, or both, indicating stromelysin synthesis immediately following resection. The extent of staining varied both between samples and within each sample, but the most extensive matrix and cellular staining occurred in rheumatoid synovia. Stromelysin 2 was never observed, but intracellular matrix-stromelysin was seen in one active inflammatory synovium. Focal synthesis of collagenase was seen in lining cells of two synovia, and in connective tissue stromal cells in four synovia. Gelatinase A was observed in stromal cells from four synovia. TIMP-1 synthesis was observed in five synovia, and in two highly active synovia the distribution of TIMP-1 positive cells was more widespread than that of MMPs. No TIMP-2 was seen in any of the synovial samples.

To our knowledge this is the first report of the occurrence of stromelysin 1 bound to extracellular matrix components of synovia. It was present in ex vivo samples from both osteoarthritic and rheumatoid synovia, but was more widespread in the latter. Matrix staining was also identified in synovial samples which contained no stromelysin secreting cells (for example those from patient 3), suggesting that the stromelysin 1 may have been bound to the matrix for some time. Synthesis of stromelysin by synovial lining and stromal cells has been reported by other authors using both immunolocalisation and in situ hybridisation methods and our data reinforce their observations that expression is focal and related to the degree of inflammation. In situ methods would not, however, detect enzymes already bound to extracellular matrix. The polyclonal antiserum used in this study recognises both the pro, active and C-terminal forms of stromelysin and all these forms bind to collagen via the C-terminal domain. Stromelysin has a broad spectrum of activity against many components of the connective tissue matrix and its presence on the matrix over prolonged periods may contribute to potentiation of the synovitis.

None of the synovia examined contained any stromelysin 2. Cultured rheumatoid synovial cells have been shown to express very low levels of stromelysin 2 mRNA, but it is likely that the six hour period of culture in monensin used for this study was insufficient to detect very small concentrations of protein synthesis. As a positive control the antiserum to stromelysin 2 was shown to immunolocalise stromelysin 2 in
NSO mouse myeloma cells transfected with stromelysin 2 (see Methods). However, matrilysin, another possible member of the stromelysin group, was identified in synovium from a patient with seronegative rheumatoid arthritis in a shoulder joint. Matrilysin was originally extracted from rat uterus, and has been shown to be a secreted product of developing human monoclonal phagocytes. This is the first report, to our knowledge, of expression of matrilysin by cells in synovium. While further work is necessary to confirm the cell type(s) secreting the matrilysin and to find further examples, it is possible that they are mononuclear phagocytes from the recent inflammatory infiltration.

Previous workers have identified collagenase in synovia by both immunolocalisation and in situ hybridisation and our results agree that collagenase may be expressed by both synovial lining and stromal cells with marked regional variations. Two rheumatoid synovia (patients 2 and 3) heavily infiltrated with inflammatory cells were found to be collagenase negative (table 2), suggesting that synthesis may be episodic as well as focal. These patients both showed active inflammatory changes in their synovium, but no significant evidence of articular cartilage erosion or degeneration. Focal synthesis of gelatinase A was observed in synovia from four patients (three rheumatoid, one osteoarthritis). It was frequently seen in connective tissue stromal cells away from the synovial surface, whereas collagenase and stromelysin were predominantly observed in the lining cells and stromal cells immediately below the lining. Focal synthesis of gelatinase A by stromal cells was also observed on days 7–28 after induction of arthritis in two rabbit models, suggesting that this enzyme has an important role in wound healing. The localisation of gelatinase A to extracellular matrix is consistent with the findings of Murphy et al, who showed that gelatinase A binds tightly to collagen through its fibronectin-like domain in addition to binding to cell membranes. Tétlow et al demonstrated synthesis of gelatinase B by a few macrophages in some synovial tissue sections. In the samples we have examined we were unable to find any gelatinase B other than in polymorphonuclear leucocytes.

All patients studied had received a variety of drug treatments before removal of synovia. It is noteworthy, however, that patient 6 had received both systemic and intra-articular steroids for several months before removal of the synovial specimen, which showed marked hyperplasia, but there was still abundant cellular synthesis of MMPs and TIMP-1. McCachren reported that two patients who had received prednisone in low dosage for a long period of time had synovia with high scores for inflammation and collagenase and TIMP mRNAs, and a recent study found no lack of stromelysin mRNA expression in two patients who had been treated with steroids. However, Firestein et al studied the effect of intra-articular corticosteroid injections on synovial tissue gene expression using serial

We thank the Medical Research Council (UK) for financial support, Mr Jim Reilly for technical assistance, and Mr Christopher Green for printing the colour photographs.


Immunolocalisation studies on six matrix metalloproteinases and their inhibitors, TIMP-1 and TIMP-2, in synovia from patients with osteo- and rheumatoid arthritis.

R M Hembry, M R Bagga, J J Reynolds and D L Hamblen

Ann Rheum Dis 1995 54: 25-32
doi: 10.1136/ard.54.1.25

Updated information and services can be found at:
http://ard.bmj.com/content/54/1/25

These include:
Email alerting service
Receive free email alerts when new articles cite this article. Sign up in the box at the top right corner of the online article.

Notes

To request permissions go to:
http://group.bmj.com/group/rights-licensing/permissions

To order reprints go to:
http://journals.bmj.com/cgi/reprintform

To subscribe to BMJ go to:
http://group.bmj.com/subscribe/