Stimulation of glycosaminoglycan production and lysosomal activity of human synovial cells in culture by low environmental pH


From the University of Melbourne, Department of Medicine, Royal Melbourne Hospital, Victoria 3050, Australia

SUMMARY Glycosaminoglycan production, acid hydrolase activity, proliferation, and morphology were examined in human synovial cells subjected to low environmental pH. The amount and the molecular size of newly synthesised glycosaminoglycan (GAG) were increased without significant change in the rate of cell proliferation. Lowered pH produced an increase in the size of cytoplasmic organelles. Some of these possessed ultrastructural features of lysosomes, but others were clearly nonlysosomal and were of uncertain identity. Intracellular activity of the lysosomal acid hydrolase N-acetyl-B-glucosaminidase (NAG) was not altered by low pH, but a marked increase occurred in extracellular NAG activity, indicating enhanced release.

Low pH values have been reported in synovial fluids from inflamed joints of patients with RA. In the most extreme cases this could enhance the activity of free lysosomal enzymes in synovial fluid, particularly for example cathespin B, which is active in the pH range of 6–7. A further possibility is that increased hydrogen ion concentrations in rheumatoid joints might induce functional changes in synovial intimal cells, which could account for a number of the pathological features of RA, including synovial proliferation, increased activity of lysosomal acid hydrolases in synovial tissue and fluid, and reduced hyaluronic acid viscosity. To examine this hypothesis glycosaminoglycan production, lysosomal activity, and proliferation of human synovial intimal cells were studied in culture conditions designed to simulate the pH changes which can occur in rheumatoid joints.

Materials and methods

Synovial cells. Primary cultures were prepared by trypsinisation of synovial tissue within intact joints of nonarthritic cadaver donors, as described previously. Cell lines derived from confluent primary cultures were used in early passages while their behaviour was still typical of euploid cells.

Accepted for publication 10 May 1983.
Correspondence to Dr B. J. Clarris.

Culture media and pH control. Eagle’s basal medium (EBM) was used throughout, supplemented with fetal bovine (FS) or human (HS) serum or both. Sera were heated 56°C for 30 min before use. Media were buffered by dilution of 1 M stock solutions of sterile NaH₂PO₄, NaHCO₃, and the organic buffers Pipes (piperazine-N,N′-bis(2-ethanesulphonic acid)), Bes (N,N-bis (2-hydroxymethyl)-2-aminoethanesulphonic acid), Hepes (N-2-hydroxyethyl-piperazine-N′-ethanesulphonic acid), Tes ((N-tris hydroxyethyl)methyl-2-aminoethanesulphonic acid), and Epps (N-2-hydroxyethyl-piperazine propane-sulphonic acid) in phosphate-free EBM as described by Ceccarini. The pH of complete media was adjusted with 1 M NaOH at 22°C, using a Radiometer (Copenhagen) 26 pH meter. pH values at 37°C were less than 0.1 pH units higher than at 22°C. Final concentrations of the individual organic buffers ranged from 5 to 20 mM. None of the buffers inhibited the growth of synovial cells at these concentrations. At the termination of experiments supernatant media were sealed in plastic tubes and the final pH was measured at 22°C.

EBM and salt solutions (Hanks’s balanced salt solutions—HBSS; Dulbecco phosphate buffered sodium chloride—PBS) were obtained from Commonwealth Serum Laboratories, and FS and trypsin from Flow Laboratories Australasia. Lipid-free bovine serum albumin (BSA) was supplied by the
Sigma Chemical Co. and the organic buffers by Grand Island Biological Co.

**Morphology and ultrastructure.** Living cultures were examined with phase contrast illumination under an Olympus PMB6 tissue culture microscope. Cultures were prepared for transmission electron microscopy (EM) by fixation in situ with 2% glutaraldehyde in 0.2 M cacodylate buffer, pH 7.4, for 10 min at room temperature, after rinsing with PBS. The cells were detached with a soft scraper, centrifuged into pellets, washed in cacodylate buffer with 7.5% sucrose, and stained with 2% osmium tetroxide for 30 min at room temperature. The pellets were dehydrated stepwise in ethanol, embedded in Epon, sectioned, and stained with lead citrate.

**Measurement of cell proliferation.** Replicate cultures were prepared in polystyrene culture flasks (25 cm² culture surface area) after dispersal of stock cultures with 0.25% trypsin in Ca²⁺/Mg²⁺-free PBS. Each flask received 4 ml EBM + 10% FS containing approximately 3 x 10⁶ cells. The vessels were sealed and the cells allowed to attach and spread at 37°C for 18-24 h. Cultures were allocated to treatments by random numbers. Before test media were added one group of cultures was selected for counting to obtain initial cell numbers. The culture vessels were sealed without adjustment of CO₂ in the atmosphere. Initial and final cell numbers were determined with a Coulter counter after complete dispersal with 0.5% trypsin in Ca²⁺/Mg²⁺-free PBS.

**Assay of NAG.** Extracellular activity was measured in 0.5 ml aliquots of the culture supernatants. After the final count 10 ml of the trypsin-cell suspension was centrifuged and the cells disrupted by freezing and thawing 3 times. The broken cells were mixed with 0.01 M tris-HCl, pH 7.0, containing 0.25 M sucrose. Intracellular NAG activity was measured in 0.5 ml of the cellular extracts after clarification by centrifugation. NAG was assayed colourimetrically as described previously.

**Measurement of glycosaminoglycan (GAG) synthesis.** Synovial cultures established in plastic flasks as described above were incubated for 48 h with [³H]-sodium acetate (specific activity 3·8 Ci per mmol; Radiochemical Centre, Amersham) at a final concentration of 20 μCi per ml (0·74 mBq per ml). The supernatants were aspirated and 0.25 ml aliquots incubated with 112 units of pronase for 60 min at 37°C. After brief centrifugation 20 μl samples were applied to Whatman 3M chromatography paper and treated with 0·1% cetylpyridinium chloride as described by Castor et al. After washing with 0·1 M NaOH [³H]-labelled GAG was measured by liquid scintillation.

**Molecular weight analysis of GAG.** Culture media containing [³H]-labelled GAG were centrifuged for 20 min at 37 000 g on a Sorvall RC2-B centrifuge. The supernatants were ultracentrifuged at 170 000 g average for 20 h at 4°C on caesium chloride gradients (initial density 1·51 g/ml). The bottom one-third of the gradient was dialysed exhaustively against 0·07 M Sørensen's phosphate buffer, pH 7·2. The molecular weight distribution of the labelled GAG was analysed by chromatography in Sephacryl S-1000 (Pharmacia Fine Chemicals), with a column 70 cm in length and 1·6 cm diameter. The column was eluted with Sørensen's buffer, pH 7·2, at a flow rate of 16·7 ml per h and radioactivity in 2 ml fractions determined by liquid scintillation counting. The elution profile was compared with molecular weight calibration curve provided by courtesy of Dr K. Granath (Pharmacia AB, Uppsala).

**Lactic acid dehydrogenase.** This was measured in culture supernatants by the Department of Biochemistry at Royal Melbourne Hospital.

**Results**

**Stability of pH.** Despite the organic buffers a drift of up to 0·3 pH units was shown by the difference between initial and final readings (Fig. 1). Single pH values shown on subsequent figures therefore represent the final reading only.

**Effect of pH on cell proliferation.** Multiplication of
Synovial cell-lines was previously found to be serum-dependent and proportional to serum concentration.\(^\text{19}\) In the present study serum concentration was standardised at 10\% (\(v/v\)). Human serum was used in preference to FS since the latter contributes a high background in the NAG assay. As shown by Fig. 1(a), a variation in pH from 7·74 to 7·08 produced no significant alteration in growth rate in 48 h. Longer exposures at pH values of approximately 7·0 usually resulted in slow loss of cells from the culture surface and littering of the medium with debris and floating cells. Further reduction in pH below 7·0 produced suppression of growth, but as shown by Fig. 1(b) some net increase in cell numbers still occurred in 2-day experiments at pH values as low as 6·8. Without serum, proliferation of synovial cells ceases, but cultures can be maintained for at least 48 h without appreciable loss of cells by replacing serum with BSA at a concentration of 4 mg per ml. As shown by Table 1, variation of pH from 7·8 to 6·6 in this medium produced no appreciable change in cell numbers over 48 h.

**GAG synthesis.** Decrease in pH produced a highly significant and reproducible increase in GAG synthesis (Fig. 2). The GAG produced at low pH was of exceptionally high molecular weight, with a narrow distribution of polymer sizes (Fig 3).

**Morphology and ultrastructure.** Macrophage-like cells present in the primary phase of isolation are strongly adherent to culture surfaces and do not readily detach with trypsin. Hence after 2-3 passages synovial cell lines consist entirely of elongated fibroblast-like cells. These normally contain many small and widely distributed cytoplasmic organelles which can be seen by phase contrast microscopy (Fig. 4A). Culture at low pH appeared to produce an increase in the size of the cytoplasmic organelles and clustering of these bodies round the nucleus (Fig. 4B), but a marked change was not evident in their number. In living cells there was no evidence of enlarged vacuoles, unusual nuclear features, or pronounced changes of cell shape in response to low pH. Typical electron microscope fields showed highly elongated cells in control cultures (Fig. 5A), whereas in cultures exposed to low pH many less elongated, more macrophage-like cells occurred, often with well developed marginal filipodia (Fig 5B). Lysosomes were sparse in the controls, and in many fields none could be seen, as in Fig. 5A. In contrast, the cells exposed to low pH contained many lysosomal bodies tending to occur in clusters. Hence the large bodies seen by phase contrast microscopy of living cells might represent lysosomal aggregates rather than grossly swollen individual lysosomes, though the mean diameter of single lysosomes was greater than

<table>
<thead>
<tr>
<th>Initial cell numbers (mean ± SEM) (\times 10^{-3}) ((n = 4))</th>
<th>Initial pH</th>
<th>Final cell numbers (mean ± SEM) (\times 10^{-3})</th>
<th>Final pH</th>
</tr>
</thead>
<tbody>
<tr>
<td>283·9 ± 3·6</td>
<td>6·50</td>
<td>292·9 ± 4·6</td>
<td>6·61</td>
</tr>
<tr>
<td>6·95</td>
<td>6·19</td>
<td>265·9 ± 11·6</td>
<td>7·19</td>
</tr>
<tr>
<td>7·37</td>
<td>7·62</td>
<td>251·5 ± 16·3</td>
<td>7·71</td>
</tr>
<tr>
<td>7·55</td>
<td>7·71</td>
<td>285·2 ± 4·4</td>
<td>7·82</td>
</tr>
<tr>
<td>7·74</td>
<td>7·82</td>
<td>288·5 ± 4·4</td>
<td>7·82</td>
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</tbody>
</table>

* Serum was replaced by BSA at 4 mg/ml.
those occurring in control cells. The phenomenon of lysosomal clustering could also explain the apparently unchanged numbers of organelles noted in the living cells. The synovial cells exposed to low pH also contained a number of amorphous, clearly non-lysosomal bodies, with approximately the same size range as the lysosomes, which remain unidentified (see example marked U in Fig. 5B).

**NAG activity.** Decrease in pH produced no significant alteration in intracellular activity of NAG in synovial cells (Fig. 6).

However, NAG levels increased in the supernatant media. NAG activity levels remained unchanged when the media were subsequently dialysed to pH 7·0 (Fig. 7), indicating that the increase was due to enhanced release from the cells rather than to any effect on buffering in the enzyme assay. Similar responses occurred in nonproliferating cultures produced by substituting BSA (4 mg per ml) for serum (Table 2).

**LDH.** None of the treatments caused appreciable release of LDH from the cells.

**Discussion**

Evidence from studies in RA1-3 suggests that tissues
within the joints in inflammatory arthritis are frequently, and perhaps chronically, subjected to high concentrations of hydrogen ions. In the present study we found that pH changes similar to those observed in rheumatoid synovial fluids can produce alterations in the behaviour of synovial intimal cells, including enhancement of GAG production and activation of the lysosomal system. The wide variations in pH alone had no appreciable effects on growth rates, and the hyperplasia commonly seen in rheumatoid synovia is therefore likely to be due to other stimuli.

Initially stimulation of GAG production by low pH was indicated by a gross and consistent increase in the amount of precipitate formed in low pH media in the 'mucin clot' test, which is used as a qualitative index of GAG production by synovial cells. Quantitative measurement by the isotopic procedure demonstrated that the differences were due largely to the comparitively high molecular weight of GAG released from the cells in conditions of low pH. Studies in this and other laboratories indicate that at least 90% of the GAG secreted by human synovial cells in culture hyaluronic acid (HA). A similar response to low pH in rat embryonic fibroblasts supports the view that increase in HA production might be a typical response of GAG-producing cells to lower pH. Our findings indicate that the low viscosity of synovial fluid in rheumatoid patients is unlikely to be a consequence of the decreased pH often noted in the joints of these patients. On the other hand the molecular weight of GAG secreted by synovial cells at pH values near to normal was considerably lower than expected from our experience with these cells cultured in standard bicarbonate-buffered medium, though variability is common in these conditions. At this stage it is not

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**Fig. 5A**

Ultrastructure of synovial cells cultured for 48 h in medium of pH (A) 7.50 (controls) and (B) 7.08 (low pH). Control cells are elongated, with sparse lysosomes (none evident in (A)). In (B) a macrophage-like cell is shown with enlarged lysosomes (L), mainly in clusters, marginal filipodia (F), and an example of an unidentified amorphous granule (U). (x 14780).

**Fig. 5B**

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cultures (+ methods'. Vertical bars are means from 4 replicate cultures (± SEM).

Table 2  Effect of variation pH in culture media on intra- and extracellular activities of NAG from synovial cells cultured in serum-free medium*

<table>
<thead>
<tr>
<th>Initial pH</th>
<th>Final pH</th>
<th>NAG†</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Intra-</td>
<td></td>
</tr>
<tr>
<td>(mean ± SEM)</td>
<td>(mean ± SEM)</td>
<td>SEM)</td>
</tr>
<tr>
<td>6-50</td>
<td>6-61</td>
<td>107-4± 8-5</td>
</tr>
<tr>
<td>6-95</td>
<td>7-19</td>
<td>135-6± 5-2</td>
</tr>
<tr>
<td>7-37</td>
<td>7-62</td>
<td>125-5± 0-7</td>
</tr>
<tr>
<td>7-55</td>
<td>7-71</td>
<td>133-8± 4-7</td>
</tr>
<tr>
<td>7-74</td>
<td>7-82</td>
<td>146-2± 2-3</td>
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</tbody>
</table>

* The medium contained BSA at 4 mg/ml in EBM. Duration of the incubation was 48 h.
† Activity of N-acetyl-B-glucosaminidase (NAG) was assayed by determining p-nitrophenol released from the substrate as described in 'Materials and methods'.

possible to specify whether the smaller molecules produced at higher pH reflect impaired polymerisation during synthesis or extracellular breakdown. Degradation due to hyaluronidase is an unsatisfactory explanation, since release of hyaluronidase from synovial cells cannot be induced, even by substances which produce gross stimulation of the lysosomal system. Another possibility is the absorption of energy from beta emission in the particular conditions of pH and buffer composition gave rise to enhanced generation of free radicals, which can cause degradation of HA. From a practical viewpoint the enhanced secretion at low pH suggests a means for optimising the production of high-grade HA for other experimental purposes.

The swollen organelles seen by phase contrast in synovial cells grown in low pH media included structures clearly identifiable by electron microscopy as lysosomes, and also bodies of about the same size as the lysosomes, but lacking recognisable features. These bodies bore a resemblance to the so-called type 1 inclusions noted by Lie et al. in human skin fibroblasts, which, however, were produced in media of unusually high pH. In both cases the accumulation of these obscure bodies in the cytoplasm might signify incipient degenerative changes, though in the case of the synovial cells such changes would not be consistent with the sustained growth rates. The apparent swelling of lysosomes in response to low pH occurred without increase in intracellular activity of NAG. However, there was a pronounced increase in extracellular NAG, and the fraction released therefore appears to be newly synthesised enzyme secreted without storage. The mechanism might involve passive loss in association with increased pinocytosis or a controlled exocytosis. This is an area for further investigation.
We have previously found that release of NAG by human synovial cells can be promoted by adenosine and also by E or F type prostaglandins, which are putative mediators of inflammation. More recently a similar reaction was found to occur in response to a soluble factor produced by human peripheral blood monocyte-macrophages (Clariss and Hamilton, unpublished results).

The total lysosomal response in the inflamed joint might therefore be due to many factors, perhaps amplified by interactions. Though the lysosomal complex contains proteolytic enzymes such as cathepsin B, which is potentially capable of acting in the breakdown of cartilage matrix, the activity of these enzymes is in general likely to be blocked after release owing to unfavourable pH, except perhaps in the most extreme cases of pH depression. On the other hand, as suggested by Dingle, much lower pH values are possible in microenvironments between closely opposed tissues at cartilage-pannus junctions. Lowered pH in these locally protected sites might induce the release and enhance the activity of lysosomal enzymes from intimal cells while partially suppressing the activity of neutral proteases such as collagenase and plasminogen activator. An analogous model for this hypothetical situation has been described by Vaes, in which degradation of matrix by osteoclasts correlates more closely with lysosomal hydrolases released into protective resorption zones than with enzymes favoured by neutral pH.

This work was supported by a grant from the National Health and Medical Research Council of Australia. The authors thank Dr K. Granath, Pharmacia AB, Uppsala, for permission to use calibration data for determination of the molecular weight of hyaluronic acid.

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B J Clarris, J R Fraser, K D Muirden, L P Malcolm, M W Holmes and K Rogers

Ann Rheum Dis 1984 43: 313-319
doi: 10.1136/ard.43.2.313

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