Cathepsin B in osteoarthritis: cytochemical and histochemical analysis of human femoral head cartilage

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Abstract

Objective—To localise the cysteine endopeptidase cathepsin B in chondrocytes and cartilage from normal and osteoarthritic (OA) human femoral heads in order to provide qualitative information on its cellular expression and distribution at possible sites of action.

Methods—OA articular cartilage was obtained at surgery for total hip replacement; control cartilage was obtained at postmortem. Chondrocytes were isolated by sequential enzymatic digestion and cathepsin B analysed by immunocytochemistry and activity staining with a fluorogenic substrate. Lysosomes were visualised by fluorescence microscopy after staining of living cells with acridine orange. Using a histochemical reaction, enzyme activity was measured in cryosections of full thickness cartilage.

Results—Chondrocytes from normal cartilage contained very few lysosomes and only a minor cell population was cathepsin B positive. A high proportion of chondrocytes from active OA cartilage contained a large number of lysosomes and an excess of cathepsin B in intracellular organelles; the enzyme was stored in an active form. In this respect, OA chondrocytes closely resembled normal cells that had been phenotypically modulated by IL-1.

Conclusion—The presence and the particular distribution of active cathepsin B in OA cartilage at 'more involved' sites suggest a pathological role for this enzyme in sustaining and perpetuating cartilage degradation. While other stimuli may also be responsible for cathepsin B expression in OA chondrocytes, the similarity with artificially modulated cells indicates fibroblastic metaplasia as a plausible mechanism.

Materials and methods

Cell isolation

OA femoral heads were obtained at total hip replacement and control specimens at postmortem. Details of sampling, documentation, definition of the anatomical position of cartilage and of its type were as in the accompanying paper; all subsequent operations were carried out under a laminar flow. The table includes a summary of the diagnoses for the OA femoral heads considered in this study. OA samples were designated by a three digit number and control specimens of cartilage identified by the prefix 'control'
using the previous numbering system, as some samples were common to both this and the previous study. The femoral heads were then washed with Hank’s balanced salt solution (HBSS) and slices of cartilage were removed with a scalpel, transferred to Petri dishes, and maintained overnight in organ culture in Ham’s F-12 medium (Gibco, Grand Island, NY, USA) containing 10% heat inactivated fetal calf serum (FCS), 0.1 mg/ml ascorbic acid (added in this high concentration only at this stage and omitted later), 100 IU/ml penicillin and 100 μg/ml streptomycin. Cartilage from OA femoral heads was sampled from osteophyte free areas of the anterior and posterior regions; control cartilage also included the superior region. The cartilage slices were then washed twice in HBSS, finely diced and the whole tissue obtained from a femoral head incubated for one hour at 37°C under gentle stirring in a mixture of 30 ml of Ham’s F-12 medium containing 54 U/ml pronase E from Streptomyces griseus (Fluka, Buchs, Switzerland) and 10 ml of 1x trypsin-EDTA solution (0.5 g/l trypsin 1:250, 0.2 g/l EDTA in modified Puck’s saline A, Gibco). The supernatant was discarded, the pellet washed twice in HBSS and digested for one to two hours with 0.2% collagenase from Clostridium histolyticum, type III, 144 U/mg (Worthington, Freehold, NJ, USA) in F-12 medium containing penicillin and streptomycin (concentrations as above) and 10% FCS. After settling of debris, the supernatants were centrifuged at 300 g and the digestion was repeated in the same way on the pellets for a further two to three hours with fresh reagents. The supernatants of the two digestions were pooled and the cells were washed twice with F-12 medium before being further processed. Cells were incubated in a humidified chamber at 37°C and an atmosphere consisting of 5% carbon dioxide in air. We demonstrated that modulation of the chondrocyte phenotype by serial monolayer subcultures in a nutrient rich medium (Dulbecco’s modified Eagle’s medium (DMEM), Gibco) resulted in a considerable increase of synthesis, storage, and secretion of cathepsin B. Within the first week in primary culture, however, cathepsin B expression was barely detectable. In the present study we did not observe stimulation of cathepsin B in primary culture for at least two weeks when the cells were cultured in a nutrient poor medium such as Ham’s F-12. Therefore, for the purpose of inducing dedifferentiation, chondrocytes from normal cartilage were plated at 2 × 10⁴ cells in 25 cm² culture flasks in DMEM supplemented with 10% FCS and 2 mmol/l glutamine. These cells were subcultured after trypsinisation of confluent monolayers. Otherwise, to evaluate chondrocytes in a condition as near as possible to that in the original tissue, the cells were plated in F-12 medium as detailed below.

**IMMUNOCYTOCHEMISTRY**

For enzyme immunolocalisation, 1 × 10⁴ chondrocytes in 0.1 ml of the appropriate medium containing 10% FCS were plated on round glass cover slips (15 mm diameter) previously sterilised with ultraviolet light. With plastic cover slips results were similar, but preparations were qualitatively inferior for microphotography. For the demonstration of the cathepsin B content in freshly isolated chondrocytes, the cells were not cultured, but simply allowed to adhere to cover slips in F-12 medium (40 hours to six days), while only the artificially modulated cells from subcultures were incubated in DMEM (see the preceding section). Adherent cells were washed with serum free medium and stained for cathepsin B with the avidin-biotin-peroxidase method using the IgG fraction of affinity purified sheep anti-rabbit cathepsin B antibodies. These cross reacted with human spleen cathepsin B purified as described. Controls were incubated with chromatographically purified non-immunised sheep IgG. Addition of FCS to F-12 medium over a short period did not alter the phenotypic traits of the cells and the rationale for using it was improvement of yield and adherence to culture vessels.

**CATEHPSIN B CYTOCHEMISTRY AND STAINING OF LYSOSOMES**

Cathepsin B activity within chondrocytes was assayed using the fluorogenic substrate N-carbobenzyloxy-l-arginyl-l-arginine-4-methoxy-β-naphthylamide (Z-Ala-Arg-Arg-MNA) (Bachem Ltd, Bubendorf, Switzerland) as the substrate, and the reaction product was coupled to 5-nitrosalicylaldehyde to reveal cathepsin B-containing organelles as bright yellow spots in the fluorescence microscope. Control reactions were performed by including either 5 μmol/l leupeptin (Bachem Ltd) or 2 μmol/l 1-trans-epoxy-succinyl-leucylamido(4-guanidino)-butane (E-64) (Sigma, St Louis, MO, USA). Lysosomes were visualised by vital staining of unfixed cells with acridine orange and fluorescence microscopy detection.

**CATEHPSIN B HISTOCHEMISTRY**

Cartilage cylinders were sampled as previously described and immediately frozen in Tissue-Tek® OCT compound (Miles, IN, USA). Only cartilage remote from the osteophytes was considered. (Osteophytes contained the greatest cathepsin B activity in an OA joint, but these results will not be discussed here.) The unfixed and undecalified blocks of tissue were allowed to reach the room temperature of ~25°C before being cryosectioned with a hardened metal knife. After the tissue block was trimmed to the desired level, a piece of transparent Scotch tape 850 (3M, St Paul, MN, USA) was fastened on to its surface using a flat, medium-hard brush. Sections 10 μm thick were obtained by cutting underneath the tape so that the tissue remained attached to it. Cathepsin B activity was visualised on the tape adherent sections with Z-Ala-Arg-Arg-MNA (Bachem Ltd) as substrate and Fast Blue BB (Serva, Heidelberg, Germany) as the post-
coupling reagent.\textsuperscript{23} \textsuperscript{24} Incubation with the substrate was in the presence of 10\% w/v polyvinyl alcohol, which very efficiently prevented diffusion of the reaction product.\textsuperscript{23} \textsuperscript{25} Control reactions were performed by including either 5 \( \mu \text{mol/l} \) leupeptin or 2 \( \mu \text{mol/l} \) E-64. The stained tissue sections adherent to tapes were mounted in glycerol and immediately photographed. The severity of the OA lesions in all samples examined was graded according to Mankin \textit{et al.}\textsuperscript{25} as detailed previously.\textsuperscript{5} Bright field and fluorescence microscopic examination, and photography of cells and tissues were performed with a Zeiss Axiophot microscope.

\section*{Results}

The table summarises details of the patients from whom OA femoral heads considered in this study were obtained. Although the primary indication for surgery of patients 007 and 011 was hip necrosis, they were included because the cartilage undoubtedly showed OA degeneration. Both femoral heads of nine individuals with no history of joint disease were analysed as controls. With OA cartilage obtained at surgery, the availability of tissue varied greatly from one sample to another, many femoral heads having just a few remnants of cartilage. Consequently, a choice was made for each as to whether cell cultures, enzyme histochemistry, or zonal enzyme distribution\textsuperscript{3} could be investigated. When only a representative few of the 1500 photographs produced in this study can be shown, the general trends they demonstrate will be considered.

\section*{CATHEPSIN B IMMUNOCYTOCHEMISTRY}

Chondrocytes from normal and OA cartilage were left in F-12 medium just for the time necessary to adhere to cover slips for immunolocalisation experiments. The incubation conditions were not sufficient for phenotypic modulation, so that cathepsin B found within the cells was likely to represent the original enzyme content. After immunostaining, cathepsin B containing chondrocytes were characterised by the presence of brown cytoplasmic granules that remained unchanged for at least one year. A diffuse, non-specific and agranular brown colour was sometimes present in the controls because of a high dosage of non-immune sheep IgG; this faded within two to four weeks in permanent mounts. Figure 1 (a–d) shows chondrocytes from four normal femoral heads (the other controls investigated behaved in the same way). Some of the cells from control cartilage adhered and spread on coverslips more easily than other cells but, independently of the donor’s age, these cells contained only a few (fig 1c+, d+) enzyme positive granules, or none at all. As they were prepared from a pool of all zones of cartilage, they were heterogeneous and the cathepsin B containing cells may represent the fibrocytic type chondrocytes from layers proximal to the cartilage surface.\textsuperscript{4}

Chondrocytes from OA cartilage contained conspicuous amounts of cathepsin B in a large number of cytoplasmic granules soon after isolation (fig 1e–h). Besides having a much greater cathepsin B content, OA chondrocytes were larger than normal cells, with a polygonal or a fibroblast like shape. To facilitate this comparison, all the photographs in figures 1 and 2 were taken using the same objective and were reproduced at the same final magnification. The OA chondrocyte population was also heterogeneous and included cells from cartilage types IV and V, but the isolation method adopted (discarding of the first pronase/trypsin digest) minimised possible contamination by foreign cells such as fibroblasts. The table summarises the cathepsin B content of the cells from nine additional patients.

**Pathological specimens obtained at total hip arthroplasty: details of origin, and analysis of cathepsin B in cells and cartilage**

<table>
<thead>
<tr>
<th>Patient, sex, age (yr), diagnosis, site</th>
<th>Cathepsin B content of cells (immunocytochemistry)</th>
<th>Cathepsin B activity in cartilage (histochemistry)</th>
</tr>
</thead>
<tbody>
<tr>
<td>001 73, PrOA, L</td>
<td>&gt;90% cells, ++++, fig 1e</td>
<td>ND</td>
</tr>
<tr>
<td>002 74, PrOA, L</td>
<td>&gt;90% cells, ++, fig 1g</td>
<td>ND</td>
</tr>
<tr>
<td>003 58, PrOA, R</td>
<td>&gt;90% cells, ++++, fig 1f</td>
<td>ND</td>
</tr>
<tr>
<td>004 55, PrOA, R</td>
<td>&gt;90% cells, +++</td>
<td>ND</td>
</tr>
<tr>
<td>005 56, Hip necrosis, L</td>
<td>~80% cells, ++</td>
<td>ND</td>
</tr>
<tr>
<td>006 50, PrOA, R</td>
<td>&gt;90% cells, +++</td>
<td>ND</td>
</tr>
<tr>
<td>007 74, PrOA, R</td>
<td>&gt;90% cells, +++</td>
<td>ND</td>
</tr>
<tr>
<td>010 73, PrOA, R</td>
<td>&gt;90% cells, +</td>
<td>P: int. in few cell clusters of mid zone, ++</td>
</tr>
<tr>
<td>011 41, Hip necrosis</td>
<td>&gt;90% cells, ++</td>
<td>P and A: int. in cell clusters of all zones, most near the surface, +</td>
</tr>
<tr>
<td>013 57, PrOA, L</td>
<td>&gt;90% cells, ++, fig 1h</td>
<td>ND</td>
</tr>
<tr>
<td>014 64, SeOA (dysplasia), L</td>
<td>ND</td>
<td>A: int. in single cells and cell clusters of all zones, ++, fig 4b</td>
</tr>
<tr>
<td>015 65, PrOA, R</td>
<td>ND</td>
<td>P: int. and ext. in all zones, ++++, fig 4e</td>
</tr>
<tr>
<td>016 57, PrOA, R</td>
<td>ND</td>
<td>A and P: int. and ext. in mid and deep zones, ++, fig 4c</td>
</tr>
<tr>
<td>017 66, PrOA, R</td>
<td>ND</td>
<td>A: int. in all zones, ++</td>
</tr>
<tr>
<td>018 66, PrOA, R</td>
<td>ND</td>
<td>A: int. and ext. around cell clusters of deep and superficial zones, ++</td>
</tr>
<tr>
<td>019 77, SeOA (traumatic), L</td>
<td>ND</td>
<td>A: int. in single cells, clusters negative, +, fig 4g</td>
</tr>
<tr>
<td>020 67, SeOA (dysplasia), R</td>
<td>ND</td>
<td>A: int. in all zones, +</td>
</tr>
<tr>
<td>021 74, PrOA, R</td>
<td>ND</td>
<td>I: int. and ext. in all zones, +++</td>
</tr>
<tr>
<td>022 53, SeOA (dysplasia), L</td>
<td>50% cells, +</td>
<td>I: int. in a few cell clusters just above the calcified zone, +</td>
</tr>
<tr>
<td>023 51, PrOA, R</td>
<td>ND</td>
<td>P: int. and ext. in all zones, ++</td>
</tr>
<tr>
<td>024 43, SeOA (dysplasia), L</td>
<td>ND</td>
<td>A: int. in all cell clusters of mid and surface zones, ++, fig 4a</td>
</tr>
<tr>
<td>025 71, PrOA, R</td>
<td>ND</td>
<td>P: int. in a few cell clusters of mid and superficial zones, +</td>
</tr>
<tr>
<td>026 50, PrOA, R</td>
<td>&gt;90% cells, +++</td>
<td>P: int. in all single cells of cartilage, but not in clusters, +</td>
</tr>
<tr>
<td>031 46, PrOA, L</td>
<td>ND</td>
<td>A and P: int. and ext. around cell clusters of mid and superficial zones, +</td>
</tr>
<tr>
<td>032 75, PrOA, R</td>
<td>ND</td>
<td>A and P: int. in all cell clusters of mid and superficial zones, +</td>
</tr>
<tr>
<td>033 57, PrOA, L</td>
<td>ND</td>
<td>A and P: int. in cell clusters of mid and superficial zones, +</td>
</tr>
<tr>
<td>034 5 85, L</td>
<td>ND</td>
<td>A: int. in single cells (+) and ext. (+) of mid-deep zones, fig 4f</td>
</tr>
<tr>
<td>036 78, PrOA, L</td>
<td>~70% cells, +</td>
<td>P: ext. in calcified zone only, +</td>
</tr>
</tbody>
</table>

Diagnoses: Pr = primary idiopathic; Se = secondary; OA = osteoarthritis. L = left; R = right. Qualitative intensity scale (histochemistry) and relative number of positive granules (immunocytochemistry): − = none; + = moderate/few; ++ = strong/large; +++ = very strong/very large; A = anterior; P = posterior; I = inferior. int. = intracellular; ext. = extracellular. ND = not determined.
PHENOTYPIC MODULATION AND STAINING OF LYSOSOMES

The cells of specimen control-5 shown in figure 1c were stained with acridine orange without prior fixation, a technique that is specific for revealing lysosomes in living cells, the lysosomes being coloured bright orange to red and the nuclei pale green. Forty hours after seeding, a large proportion of the cells did not contain appreciable amounts of lysosomes, some cells contained a discrete number of small, acridine orange positive granules, and only a very few contained lysosomes in sizeable amounts (fig 2a). The same chondrocytes (control-5) were subcultured in DMEM in the presence of 10% FCS. Figure 2b shows that these cells were considerably enlarged and contained dramatically more cathepsin B positive granules after the third subculture, as detected by immunocytochemistry. The cells of other controls behaved identically (not shown). Also, the number of lysosomes stainable by acridine orange increased considerably after the third subculture in DMEM (fig 2c). Chondrocytes from OA cartilage (not modulated by subculture) contained a large number of lysosomes, comparable to that seen in artificially modulated cells (fig 2d, e).

CATHEPSIN B CYTOCHEMISTRY IN OA CHONDROCYTES

From the results in the preceding sections (compare figure 1f,g and 2d,e) it is evident that the number of lysosomes in OA chondrocytes is much greater than the cathepsin B positive

Figure 1  Immunolocalisation of cathepsin B in human femoral head chondrocytes. a: Control-2, 37 yr; b: control-4, 25 yr; c: control-5, 30 yr; d: control-8, 78 yr; e: patient 001; f: patient 003; g: patient 002; h: patient 013 (patients' data summarised in the table). + = Staining with sheep antibodies against cathepsin B; - = control with non-immune sheep IgG. The 20 μm bar shown in panel a is valid for all other panels.
granules as detected by immunocytochemistry. A further method was thus applied to stain cell organelles for cathepsin B activity (for example figure 2f), and this confirmed that not all of the lysosomes made visible by acridine orange do contain cathepsin B.

CATHEPSIN B HISTOCHEMISTRY

Unfixed and undecalcified cryostat sections of normal and OA articular cartilage were stained for cathepsin B activity using a specific and sensitive method (figure 3 and 4; all photographs oriented with the cartilage surface up, cathepsin B containing areas showing as various shades of red). Two to four plugs of normal cartilage from each femoral head of nine individuals were carefully examined at high magnification in slices from various parts of the plugs. None of the specimens of control cartilage was stained positively for cathepsin B, as judged by comparing stained sections in the presence and absence of E-64. Three examples, from individuals aged 30, 54 and 78 years are shown in figure 3a–d. To save space, the control reaction, performed in the presence of E-64 as cathepsin B inhibitor, is shown for just one sample (fig 3b). Cathepsin B activity
Figure 3  Histochemical demonstration of cathepsin B in unfixed, undecalcified cryostat sections of human femoral head cartilage.  
a, b: Control-9, 54 yr (a1 and b1 lower zones, a2 and b2 upper zones of cartilage);  
c1 and c2: control-5, 30 yr, lower and upper zones, respectively;  
d1 and d2: control-8, 78 yr, lower and upper zones, respectively;  
e: patient 027 (data in table) (e1 deep zone, e2 mid zone, e3 exposed, fibrillated surface).  
+ = Positive staining; - = control in the presence of E-64. All bars represent 50 μm.

was not found in apparently intact or slightly damaged OA cartilage, while areas showing clear signs of destruction or remodelling stained positively (figs 3e, 4a–g). In all but one of the positive specimens (patient 036 in the table), enzyme activity was found within chondrocytes, while in just eight cases cathepsin B was also present in the matrix, and in patient 036 it was present only in the matrix (table). The assay solution included 10% polyvinyl alcohol to prevent enzyme diffusion and its omission caused colour diffusion as a gradient within a short range from the cells. Since activity in the matrix was not general to all
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Figure 4  Histochemical demonstration of cathepsin B in unfixed, undecalcified cryostat sections of human femoral head cartilage.  

<table>
<thead>
<tr>
<th></th>
<th>a1+</th>
<th>a2+</th>
<th>a3-</th>
<th>b+</th>
</tr>
</thead>
<tbody>
<tr>
<td>a1</td>
<td>Patient 028, mid zone and near the exposed, fibrillated surface</td>
<td>Patient 014, mid zone</td>
<td>Patient 016, chondrocytes from just above the calcified zone, c2 and c3 from the mid zone</td>
<td>Patient 032, mid zone</td>
</tr>
<tr>
<td>c1</td>
<td>Patient 015, the remaining cartilaginous layer was covered by fibrous pannus (only cartilage shown)</td>
<td>Patient 035, mid-deep zones</td>
<td>Patient 019, mid zone</td>
<td></td>
</tr>
</tbody>
</table>

Patients' data summarised in the table. + = Positive staining; - = control in the presence of E-64. All bars represent 50 μm except in panels c1–c3 (10 μm).

specimens and sections within the same specimen, and we adopted the necessary technical precautions, we describe this activity as being extracellular. In the cartilages of at least two patients (figs 3e, 4e), extracellular activity was a regular feature in all sections of all plugs examined, and we trust that this finding is not an artefact of diffusion.

The cartilage sections in figure 3e were from a patient with primary OA (Mankin score 7–9 at that site) and show intracellular and extracellular cathepsin B activity in the deep
zona (fig 3e1), in the mid zone (fig 3e2), and at the fibrillated surface (fig 3e3). Figure 4a shows the cartilage from a patient with secondary OA (hip dysplasia, Mankin score 11-12). Cathepsin B was present in clustered chondrocytes in the mid zone, near the exposed surface and along clefts penetrating into the depth, while it was absent in the deep zones. Figure 4c (Mankin score 7-8 for these slices) shows cathepsin B just above the calcified zone (fig 4c1) and in the mid zone (fig 4c2; control reaction in fig 4c3). Figure 4e illustrates a cartilage with high cathepsin B activity from a patient with severely erosive OA, conspicuous synovial inflammation and covering of the residual cartilage by fibrous connective tissue pannus. The Mankin score of this sample was 8-9, and enzyme activity was more intense towards the surface in contact with inflamed tissue, thus suggesting its participation in cathepsin B expression. This case was the only one of 36 investigated in this and the parallel study 6 which showed a conspicuous pannus invasion of remaining OA cartilage. In the OA cartilage shown in figure 4f, cathepsin B positive chondrocytes were observed in the mid and deep zones, while the upper zones were negative (Mankin score 4-5). Figure 4b, d and g show cathepsin B activity in chondrocytes from three OA cartilages. Data pertinent to these patients, and additional data, are summarised in the table, in which the intensity of the cathepsin B activity reaction is qualitatively evaluated. Many cathepsin B negative sections were found adjacent to positively stained sections within a cartilage plug, and there was no activity in some plugs adjacent to highly positive samples. The table includes data for only one OA cartilage that was cathepsin B negative in all plugs and sections examined (patient 020), for reasons of space. As it was impossible to examine all sections obtainable from a plug and we did not have a quantitative mean for defining histochemically determined cathepsin B activity, we did not attempt a statistical evaluation of the data.

Discussion

Cathepsin B activity was not found in normal cartilage using a histochemical method, but the enzyme was seen in OA cartilage at sites involved in matrix destruction and repair. The present results agree with and complement the study of zonal distribution of enzyme activity determined by a quantitative method. 6 They also extend valuable previous work on this subject 1-4 by showing that the enzyme of enzyme activity in relation to anatomy and degree of joint cartilage destruction. Active cathepsin B was found within cells, particularly in clusters of proliferated cells, and more rarely in the extracellular space. There is an apparent discrepancy between the absence of histochemically determined cathepsin B in normal cartilage and the low, but still measurable, zonal activity measured with a fluorogenic substrate. 6 For histochemistry, the cartilage slices were carefully protected to prevent cell disruption, so that normal chondrocytes, that contain just a few granules with cathepsin B, may not be accessible to the substrate because of the natural barriers constituted by the cell wall and the mucous nature of the cell surface. They may have the cathepsins masked by the presence of naturally occurring inhibitors, or may have enzyme levels below the limit of detection of the method. For the measurement of zonal enzyme distribution, cryosectioned slices were thawed rapidly in the assay buffer in order to disrupt the cells and detect all the activity of the slices. 6

Chondrocytes isolated from OA articular cartilage had some morphological and biochemical properties observed in chondrocytes from normal cartilage that were phenotypically modulated by serial subcultures in monolayers. Characteristic features of both OA and artificially modulated cells were larger dimensions and an increased number of lysosomes. A subpopulation of the lysosomes contained immunospecifically, enzymatically active cathepsin B. A disadvantage of the methods used for cathepsin B and lysosome staining in chondrocytes is that we cannot demonstrate whether all or just a subpopulation of the cells adhered to the glass cover slips. Differences in adherence were clearly seen between normal and OA chondrocytes. However, OA cells that adhered to and spread on cover slips had larger dimensions than controls cells. A constant feature was that control chondrocytes, even if left to attach and spread for several days in F-12 medium, never contained the large numbers of lysosomes and of cathepsin B positive granules observed in OA cells. The relevant result, even if qualitative, is that chondrocytes have a decidedly increased proteolytic potential are present in OA cartilage.

Previous ultrastructural studies offered convincing evidence that chondrocytes, in the advanced phases of OA, undergo progressive enlargement, contain many more intracellular organelles, and are capable of replication. 26,27 This indicates a potentiation of their synthetic and secretory activities in a way that can be mimicked in vitro through serial subcultures in monolayer. Analogous morphological changes have also been described in pig articular cartilage cultivated in the presence of retinol 28 and in co-cultures of pig cartilage with synovium or invading marrow in the presence of complement-sufficient rabbit antisera and pig erythrocytes. 29,30 With this information in mind, if we accept our previous proposal that cathepsin B is a characteristic product of phenotypically modulated (dedifferentiated) chondrocytes, we propose the working hypothesis that the expression of cathepsin B in OA cartilage may be a product of chondrocytes that have undergone 'fibroblastic metaplasia'. This hypothesis does not preclude the (co)existence of other mechanisms. The high enzymatic content of these cells probably reflects a physiological demand for increased turnover of proteins in a tissue subject to remoulding, but will any activity leaking into the matrix be sufficiently controlled? The pooled results on histological, histochemical, immunocytochemical (this
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