Expression of CD44 in normal and rheumatoid synovium and cultured synovial fibroblasts

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Abstract

Objective—To determine if expression of CD44, the principal receptor for hyaluronan, was altered in rheumatoid (RA) synovium and cultured rheumatoid synovial fibroblasts.

Methods—Synovium was obtained from normal adult human joints (n = 4) and from joints of patients with RA (n = 5). Specific monoclonal antibodies to CD44 were used in immunofluorescence of whole synovium and cultured synovial fibroblasts and in quantitative Western blotting and ELISA of CD44 in cultured synovial fibroblasts.

Results—CD44 was restricted to the lining layer in normal synovium but present, in reduced concentrations, throughout rheumatoid synovium. Cultured rheumatoid cells were 19% larger in area and showed far fewer and less extensive CD44-positive cytoplasmic extensions, together with reduced staining intensity compared with normal. Quantitative Western blotting normalised for cell protein showed a 75% reduction (normal = 1754 (835), rheumatoid = 409 (84) mean (SD) arbitrary units) in the amount of CD44 in rheumatoid cells compared with normal, and enzyme linked immunosorbent assay (ELISA) of cultured cell monolayers normalised for cell number indicated a 29% reduction (normal = 0.707 (0.110), rheumatoid = 0.504 (0.103), mean (SD) optical density at 405 nm).

Conclusions—Rheumatoid synovial cells showed altered morphology and reduced CD44 expression compared with normal cells. CD44, by means of modulated associations with the cytoskeleton, may be involved in cell shape change.


Materials and methods

MONOCLONAL ANTIBODIES

Table 1 gives details of the monoclonal antibodies (MAb) used. Antibodies were used at an appropriate dilution optimised by titration for each technique. The three antibodies to CD44 used in this study recognise the hyaluronan binding form of CD44: F.10.44.2 and A3D8 bind to the same determinant, whereas F.10.62.1 binds to a separate site on the CD44 molecule.
Tissues

Normal synovium (n = 4) was obtained from the knee joints of patients undergoing amputation for localized proximal sarcomata. Tissue was taken from macroscopically normal sites at least 30 cm distal to the tumour. Diseased synovium (n = 5) was obtained from patients with definite or classic rheumatoid arthritis according to the criteria of the American Rheumatism Association,24 undergoing major joint surgery. Mean age of control tissue donors was 63.5 years and that of RA patients, 68 (difference not significant by Student’s t test at the 95% level). All tissues were processed within two hours of removal and trimmed of capsule and fat. Blocks of approximately 5 mm³ were snap frozen in a bath of isopentane (British Drug Houses, UK) and solid carbon dioxide and stored in air tight vials at −70°C. Sections were cut at a thickness of 7 μm on a cryostat, dried under a cool fan, wrapped in tin foil and stored at −70°C.

Cell Culture

Experiments were performed with cells which were harvested before the third passage and all cultures were supplemented with antibiotics. Cells were obtained from synovial tissue by three methods.

Explant technique—This was used for both normal and rheumatoid tissue. Blocks of fresh synovium of size 1 mm³ were placed intimal side down onto the surface of T25 tissue culture flasks which had been prewetted with Dulbecco’s minimal essential medium (DMEM) containing 10% fetal calf serum (FCS) (Tissue Culture Services, UK). After 30 minutes at 37°C the explants were adherent and 5 ml of DMEM containing 10% FCS was added to each flask. The flasks were then left at 37°C in a humidified incubator under 5% carbon dioxide until a substantial outgrowth of cells was apparent (approximately seven days). The explants were then removed with forceps and the cells detached by 2-5% trypsin in phosphate buffered saline (PBS), replated in fresh flasks and grown to confluence before further passage.

Trypsin digestion of the synovial lining—Used for normal synovium. The synovial surface was washed with PBS before being treated with 2.5% trypsin solution for one hour at 37°C. The synovial surface was then gently scraped and the resulting cell suspension transferred in volumes of approximately 0.1 ml to the wells of a 96 well tissue culture plate. The cells in each well were grown to confluence before being serially cultured through intermediate sized culture plates into T25 flasks.

Collagenase digestion—This method was used for the disaggregation of RA tissue in which the distinction between the synovial surface and the deeper tissue was less clear. Synovium was chopped into small pieces and added to a solution of collagenase ( Worthington Biochemical Corp., USA) at 2 mg/ml in DMEM containing 10% FCS. After incubation at 37°C for two hours, the suspension was filtered through gauze, the cells centrifuged into a pellet at 120 g for 10 minutes and seeded at 10⁶ cells/ml in T25 flasks.

Non-specific esterase activity

Non-specific esterase activity was estimated using a standard α-naphthyl acetate technique.25

Measurement of cell surface area

Cells in 2% FCS were seeded at subconfluence onto cover slips and incubated overnight at 37°C. Cover slips were then washed in PBS before the cells were stained with 0.1% toluidine blue (Sigma, UK) in 0.1 mol/l acetate buffer, pH 5.2. Cells were rinsed with distilled water and air dried. Cell areas were measured using a Zeiss standard microscope fitted with a Panasonic WVBL 600 video camera connected to an Apple Macintosh IIfx computer running NIH image software.

Indirect immunofluorescence

For immunofluorescence studies, cells in DMEM containing 2% FCS were seeded at subconfluence onto cover slips and incubated overnight at 37°C. Cover slips were then washed in PBS before the cells were stained with 0.1% toluidine blue (Sigma, UK) in 0.1 mol/l acetate buffer, pH 5.2. Cells were rinsed with distilled water and air dried. Cell areas were measured using a Zeiss standard microscope fitted with a Panasonic WVBL 600 video camera connected to an Apple Macintosh IIfx computer running NIH image software.

Quantitative Western blot analysis

Cell monolayers were detached from culture plates by incubation with 16 mmol/l EDTA in PBS for 20 minutes at 37°C and centrifuged for 10 minutes at 1000 g. The cell pellet was solubilised in Triton buffer (300 mmol/l NaCl, 1% Triton X-100, 50 mmol/l Tris pH 7.5, 2 mmol/l CaCl₂, 2 mmol/l DTT). The solubilised samples were subjected to 15% SDS-PAGE and transferred to nitrocellulose membranes. Membranes were blocked with 10% normal goat serum (Sigma, UK) and probed with the appropriate antibodies as described in Table 1.
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50 mmol/l Tris HCl pH 7·6, 0·5% Triton X-100, 2 mmol/l phenyl methylsulphonyl fluoride) for 45 minutes on ice and the lysate centrifuged to remove cell nuclei. Ten percent sodium deoxycholate and 10% sodium dodecyl sulphate were added to the supernatant to a final concentration of 0·2%. Protein content was estimated by bicinchoninic acid (BCA) assay (Pierce, UK), and samples of supernatant of equal protein content were electrophoresed on a 7·5%-polyacrylamide gel under non-reducing conditions. Proteins were transferred to nitrocellulose membranes and duplicate blots were stained for CD44 using MAbs F.10.44.2, F.10.62.1, and A3D8. The secondary antibody was an alkaline-phosphatase-labelled goat antimouse Ig (Southern Biotechnology, USA). Binding was visualised with the colour development substrates 5-bromo-4-chloro-3 indolyl phosphate and nitroblue tetrazolium (Sigma, UK). Intensity of staining of the CD44 bands was measured by densitometry. Blots were scanned using a Hewlett Packard Scanjet 2C scanner connected to an Apple Macintosh IIx running NIH Image software with a gel plotting macro.

CELL BASED ELISA

Cellular CD44 expression was measured using a cell monolayer ELISA based on the method of Chin et al. Cells were seeded in DMEM containing 2% FCS at a density of 3 x 10^5 cells/well in 96 well tissue culture plates and allowed to adhere before being incubated overnight with 0·9%, 2%, or 10% FCS. The medium was removed and the cell layers washed twice with Tris buffered saline (TBS), pH 7·6, before fixation in 2% formaldehyde in TBS for five minutes. After a further wash in TBS, the cells were incubated with binding buffer (0·1% bovine serum albumin in TBS) for 60 minutes at 37°C. The binding buffer was replaced by the primary antibody in binding buffer and incubated for 45 minutes at 37°C. Cells were washed twice with TBS and incubated with binding buffer for 60 minutes at 37°C. The binding buffer was then replaced with alkaline-phosphatase-conjugated goat anti-mouse IgG (Southern Biotechnology, USA) at 0·2 μg/ml in binding buffer and the cells incubated for 45 minutes at 37°C. After washing in TBS, cells were incubated with Sigma 104 phosphatase substrate (Sigma, UK) dissolved in bicarbonate buffer (7·5 mmol/l Na2CO3, 18 mmol/l NaHCO3, pH 9·6) for 60 minutes. The reaction was terminated by the addition of 25 μl of 12·5 mol/l sodium hydroxide. One hundred microtitre aliquots of the supernatant were transferred to 96 well microtitre plates and the absorbance at 405 nm determined using a Dynatech MR4000 ELISA plate reader. Standard curves were established for each cell line and all assays were performed within the linear region of each standard curve.

STATISTICAL METHODS

Measurements of cell surface area were performed on samples of 40 cells per cell line, ELISAs were performed in septuplicate and results processed using Student’s two tail t test. Western blot results were expressed as means of values from the two groups (normal and RA) and the difference between them tested by Student’s two tail t test.

Results

The findings were consistent between different specimens within the normal and rheumatoid groups and did not appear to show significant variation with patient characteristics such as age, sex, disease duration, and antirheumatic drug therapy.

CULTURED CELLS

Cells obtained by explant and enzyme digestion from the same tissue source exhibited no differences in appearance, growth rate or CD44 expression. Cells cultured from synovial tissue had the general appearance of fibroblasts, but it is known that enzymatic digestion of synovial tissue releases large numbers of macrophages as well as fibroblasts, although the macrophages seem not to survive beyond the first passage. In the absence of a specific fibroblast marker, determination of the fibroblastic nature of the cells was aided by positive staining for prolyl hydroxylase, absence of esterase activity and absence of high affinity Fcy receptors (data not shown).

Cultured RA synovial fibroblasts were larger (mean = 2226 arbitrary units of area) than normal cells (mean = 1866 arbitrary units of area). This difference was found to be statistically significant (p < 0·005).

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In normal synovium, the bodies of the lining cells and sparsely scattered subsynovial cells were densely stained for CD44 (fig 1A). The intervening matrix showed many densely stained fibres which appeared to be cellular processes. These were closely packed in the lining layer but more widely separated in the deeper layers where the cells were scattered. The overall appearance was of dense staining throughout the lining layer, with minimal staining of the subsynovium. Rheumatoid synovium (fig 1B), in contrast, showed staining of almost all cells, including lymphocytes and vascular smooth muscle cells, throughout the depth of the tissue. The staining of the lining layer was markedly less dense than that seen in normal synovium. In both normal and diseased tissue, endothelial cells were uniformly unstained.

The findings in cultured synovial fibroblasts were consistent with but less marked than those in whole tissue. Normal fibroblasts (fig 2A) were uniformly stained and showed numerous multiply branched CD44-positive cytoplasmic processes which appeared to connect with similar processes from neighbouring cells. Rheumatoid fibroblasts (fig 2B) were less intensely stained for CD44 and showed far fewer processes and cellular interconnections.
Results shown are for MAb F.10.44.2, but similar results were obtained with each of the antibodies to CD44. Cell lyase preparations were normalised for total cellular protein, so it would appear that RA fibroblasts contain less CD44 than normal for an equivalent mass of cell protein.

**CELL-BASED ELISA**

There was no significant difference in optical density values obtained with normal and RA cells when they had been incubated with 2% and 10% FCS. However, at the (more physiological) concentration of 0.5%, a difference of 29% was observed (p < 0.001) (table 2).

Results shown are for MAb F.10.44.2, but similar results were obtained with each of the antibodies to CD44.

**Discussion**

This study has shown by a number of methods that CD44 is expressed by synovial fibroblasts and that its expression may be reduced in rheumatoid cells.

In normal synovium, CD44 was present predominantly in the lining layer, an area known to be largely populated by fibroblasts, where its presence on prolonged cytoplasmic processes which minutely subdivide the matrix may help in anchorage and matrix turnover. In contrast, CD44 was present on the majority of cells, albeit expressed to a lesser extent, throughout the depth of rheumatoid synovium.

Morphological studies show hyaluronan to be distributed predominantly to the lining layer in normal synovium but to be present uniformly throughout the depth of rheumatoid tissue, and this study shows that synovial distribution of the hyaluronan receptor, CD44, mirrors that of hyaluronan. The finding that CD44 and its principal ligand, hyaluronan, are localised primarily to the intima in normal synovium and that no such restricted distribution is present in rheumatoid tissue adds to a number of previously identified features of the normal synovial lining which are altered in rheumatoid arthritis. These include activity of the enzyme uridine diphosphoglucose dehydrogenase (important in the synthesis of hyaluronan precursors) and expression of chondroitin-6-sulphate, vascular cell adhesion molecule-1, and type VI collagen.

A recent study by Johnson et al found that, for a number of cell types, the percentage of CD44-positive cells was increased in RA synovium compared with normal and it was concluded that CD44 is upregulated in RA. However, individual cell types were identified on serial sections rather than by double labelling and no attempt was made to quantify the level of expression of CD44. Certainly, in RA synovium there is a massive influx of cells (largely macrophages and lymphocytes) which constitutively express CD44, but we have found no evidence for upregulation of CD44 on individual cells.
Figure 2 CD44 staining and normal (A) and RA (B) synovial fibroblasts. Normal cells show intense staining and many cytoplasmic processes. RA cells, in comparison, show reduced staining intensity and far fewer cytoplasmic processes. Bar = 10 μm.

In another study by Haynes et al., purporting to show that CD44 expression was upregulated in rheumatoid compared with normal synovium, equivalent weights of homogenised tissue were subjected to Western blotting and probed for CD44 content. Given, however, that CD44 appears to be purely cell associated, such studies require control for differences in tissue cellularity. An increase in tissue cellularity may overwhelm a concomitant decrease in individual cellular expression, resulting in an overall increase in tissue total CD44 content.

Further investigation of CD44 in the present study was performed on pure populations of cultured synovial fibroblasts at early passage. Relevant to this study is the fact that, in vivo, rheumatoid synovial cells are exposed to interleukin-1β, interleukin-6, and tumour necrosis factor α. These cytokines have been shown to modulate the expression of extracellular matrix and cell adhesion molecules and it should be noted that their lack in culture may influence CD44 expression.

Western blot analysis, normalised for cell protein, and immunofluorescent staining each suggested that there was a decrease in CD44 expression in cultured rheumatoid cells. Cell based ELISA also showed a reduction in CD44 expression per cell in RA compared with normal, but only when the concentration of FCS was decreased to 0.5%. For synovial cells serum is not the physiological fluid which they encounter in the synovium and the difference in CD44 expression between normal and RA cells may be reduced in the presence of mitogenic factors in the greater concentrations of serum. Nevertheless, the difference in CD44 expression in the presence of 2% serum was still of sufficient magnitude to be detected by Western blot analysis and immunofluorescence. However, in the cell based ELISA, differences in cell size could be a confounding factor. We have shown that cultured RA synovial cells are larger than normal and this may partially mask a reduction in receptors per unit surface area when CD44 expression per cell is measured in the cell based ELISA.

Immunofluorescent staining of cultured synovial fibroblasts showed that normal cells have CD44-positive processes which are not present on rheumatoid cells, indicating an alteration in the cytoskeleton. It is known that CD44 interacts with the cytoskeleton. A study using a mouse macrophage model of inflammation found a difference in CD44 phosphorylation and cytoskeletal association between resident cells and those which had been elicited to migrate to a site of inflammation. An earlier report found that cytoskeletal elements in rheumatoid synovial lining cells differed from those in normal cells. We have demonstrated a difference in cell shape and area between normal and RA cells and this supports the view that the cytoskeleton is altered in RA cells. We conclude that the role
of CD44 in RA may involve modulation of molecular associations of CD44 within the cell, together with a decrease in the total amount of cellular CD44.

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