Localisation of collagen types and fibronectin in cartilage by immunofluorescence

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SUMMARY Collagens type I, II, III, IV, and V and the minor cartilage collagens, 1α 2α 3α, C-PS 1, and C-PS 2, were purified, antibodies raised, and then used in immunofluorescence studies on bovine nasal cartilage (BNC). Punctate localisation was seen with the type II antibody. However, pretreatment of sections with hyaluronidase to remove the proteoglycans resulted in diffuse staining over all the section with this antibody. Antibodies to 1α 2α 3α, C-PS 1, and C-PS 2 collagens gave no staining on untreated BNC sections, but after treatment with hyaluronidase all 3 antibodies showed as a diffuse ‘halo’ round each chondrocyte lacuna. Anti-type I, anti-type III, and anti-type IV collagen antibodies did not stain untreated or enzyme treated BNC. Type V collagen antibodies gave a bright ring in the pericellular region of the lacunae of hyaluronidase-treated BNC. This was unexpected, as we could not detect type V collagen biochemically in the same cartilage. Anti-fibronectin antibodies stained areas distant from the chondrocytes, these areas being distinct from those stained by 1α 2α 3α and C-PS antibodies, suggesting that fibronectin is not associated with these collagens in BNC. These results suggest that different collagen types may have different locations within the cartilage matrix, that proteoglycans may inhibit antibody association with collagen, and that fibronectin is normally not associated with all types of collagen.

At least 5 structurally and genetically distinct and well defined types of collagen have been identified. The predominant collagen of cartilage is type II, which has been localised in the cartilage matrix by specific antibodies. Proteoglycans of human and bovine hyaline cartilage have been purified and localised by immunofluorescent techniques.

The presence in cartilage of small quantities of other collagenous components has been reported. Type V collagen, consisting of α1 (V) (previously called αB) chains only, has been described in epiphysial cartilage and shown by immunofluorescence to be located pericellularly around the chondrocyte. Burgeson et al. isolated and partially characterised 3 α-sized collagen chains, designated 1α, 2α, 3α, from human hyaline cartilage. The 1α and 2α chains are similar to the α chains of type V collagen but are genetically distinct. The 3α chain, although virtually identical with the α1 (II) chains of type II collagen, is more glycosylated and consistently copurifies with the other 2 chains. The collagens of bovine nasal cartilage (BNC) have been studied by Ayad et al., who identified type II and 1α 2α 3α collagens and 2 new collagens designated C-PS 1 and C-PS 2. C-PS 1 and C-PS 2 collagens are short triple helical collagens approximately one-third and one-seventh the length of type II collagen respectively and the constituent chains are linked by interchain disulphide bonds.

This report describes the localisation of these collagens in BNC by immunofluorescence.

Materials and methods

Collagens. These were isolated from various bovine tissues after limited pepsin digestion. Type II, 1α 2α 3α, C-PS 1, and C-PS 2 collagens were isolated from adult nasal cartilage as described by Ayad et al. Types I and III collagen were prepared from placental membranes according to Epstein. Types IV and V collagen were isolated from placental villi and membrane respectively by the method of Abedin et
al. Each collagen was purified by repeated precipitation according to its specific solubility properties in 0.02 M NaOH, HPO₄²⁻ (pH 9.2), phosphate-buffered saline (PBS) (pH 7.2) and 0.5-0.5 M NaCl. 0.05 M tris HCl (pH 7.4). The purity of these preparations was checked by sodium dodecyl sulphate polyacrylamide gel electrophoresis.

Antisera to collagen. Antisera to bovine native type II collagen were raised in rats injected intradermally with an emulsion of type II collagen in incomplete Freund's adjuvant (ICFA, Miles), by the regimen of Morgan et al. A proportion of rats developed type II collagen-induced arthritis, and separate pools of sera from arthritic and nonarthritic rats were made. IgG antibodies to native type II collagen in each pool were measured by radioimmunoassay, and were 87-1 µg of labelled anti-rat IgG bound/ml of serum for the arthritic pool and 59-6 µg for the nonarthritic pool.

Antiser to bovine native types I-V, 1α 2α 3α, C-PS 1, and C-PS 2 collagens were raised in guinea-pigs by an immunisation regimen similar to that of Beard et al. Guinea-pigs were inoculated intradermally with 500 µg of collagen in complete Freund's adjuvant (Difco) and boosted intradermally 2-3 weeks later with 250 µg collagen in ICFA (Difco). Two weeks after the boost animals were bled out by cardiac puncture under halothane anaesthesia.

Immunoabsorption of antiser. 1α 2α 3α, C-PS 1, C-PS 2, types I, II, III, IV, and V collagen were coupled to cyanogen-bromide-activated sepharose (Pharmacia), by a modification, described in the manufacturer's literature, of the method of Axen et al. Each antiserum was purified by absorption against the homologous collagens and finally absorbed on to a column of the homologous collagen. Specific antibodies were eluted with glycine-HCl buffer, 0.2 M, pH 2.2, neutralised, freeze-dried, and reconstituted in PBS at 1 mg/ml. Antibodies were routinely diluted 1/10 and 1/20 before use.

Antiserum to fibronectin. Antiserum raised in rabbits against human fibronectin was obtained commercially (Miles Laboratories Ltd). The product was an IgG fraction of the antiserum and gave identical results to a fibronectin antiserum, also used, which was raised as described by Scott et al.

Sectioning of BNC and enzyme treatments. Blocks of BNC were snap-frozen in n-isopentane cooled by liquid nitrogen and stored at -70°C. Sections 5 µm thick were cut on a cryostat and transferred to microscope slides coated with a solution of 2% Bostik clear adhesive (Bostik Ltd, Leicester) in acetone. Pretreatment of slides in this way assisted adherence of sections. To expose collagen by removal of the proteoglycans, sections were treated with hyaluronidase (Worthington Biochemical Corporation, New Jersey), diluted to 186 units/ml in 0.025 M NaCl/0.05 M acetate buffer, pH 5.0. Sections were incubated with the enzyme for 30 minutes at room temperature, then washed in PBS before staining. Some other sections were incubated with bacterial collagenase R (Calbiochem, San Diego, California), at 38 units/ml for 30 minutes at room temperature and washed with PBS before use.

Alcian blue and Sirius red stains. Sections were stained for proteoglycan with alcian blue, pH 2.5, by the method of Lison and for collagen with Sirius red by the method of Sweat et al.

Immunofluorescence. Sections were incubated with collagen antiser for 30 minutes washed with 3 changes of PBS (30 min) and then stained with either FITC-conjugated goat anti-guinea-pig immunoglobulins, FITC-conjugated rabbit anti-rat immunoglobulins, or FITC-conjugated swine anti-rabbit immunoglobulins (Nordic Immunological Laboratories Ltd, Maidenhead, Berks), as appropriate. After further washing in PBS, sections were mounted in glycerol/saline and observed with a Vickers M41 Photoplan microscope, fitted with a xenon lamp. Controls included substituting normal rat serum, normal guinea-pig serum, and PBS for anti-collagen antibodies in the first stage and blocking the fluorescent staining with unlabelled anti-rat or anti-guinea-pig immunoglobulins.

Results

Description of cartilage.

Regions of the cartilage matrix are described according to Poole et al. (Fig. 1). The zone at the surface of the chondrocyte and at the edge of the lacuna is referred to as the pericellular area. The wider zone surrounding this is the territorial zone, and the region between the territorial zones is the interterritorial matrix.

![Fig. 1 Areas of the cartilage matrix. The zone at the surface of the chondrocyte is referred to as the pericellular area. The wider zone surrounding this is the territorial zone, and the region between territorial zones is the interterritorial matrix.](http://ard.bmj.com/Downloaded from)
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Fig. 2 BNC stained with Sirius red. (×470).

Fig. 3 BNC treated with hyaluronidase, then stained with Sirius red. (×470).

Fig. 4 BNC stained with alcian blue. (×470).

Fig. 5 BNC treated with hyaluronidase, then stained with alcian blue. (×470).

Fig. 6 Immunolocalisation of type II collagen on untreated BNC. (×470).

HISTOCHEMICAL STAINING OF BNC

A section of untreated BNC stained with Sirius red is shown in Fig. 2. Fig. 3 shows a section of BNC treated with hyaluronidase and stained with Sirius red. Hyaluronidase treatment resulted in increased, more dense staining, especially in the territorial zone, indicating that more collagen was exposed. Figs 4 and 5 show BNC stained with alcian blue, untreated and after hyaluronidase treatment respectively. Decreased staining after treatment indicates that proteoglycan has been removed by the enzyme. Hyaluronidase treatment frequently resulted in displacement of chondrocytes, which were washed away during the staining procedures, as the sections were not fixed.
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Type II collagen. Immunolocalisation of type II collagen on untreated BNC gave discrete groups of fluorescent spots in the matrix (Fig. 6). Chondrocytes stained brightly, but, as this was also seen in controls with normal rat serum and unconjugated blocking antibodies (Figs 7 and 8), it was considered to be nonspecific. Diffuse staining over the cartilage matrix was observed on hyaluronidase-treated sections (Fig. 9), which was absent in controls. Qualitatively similar staining patterns were observed with sera from arthritic and nonarthritic rats.

Staining with anti-type II collagen antisera was abolished by treatment of sections with bacterial collagenase (Fig. 10).

1α 2α 3α collagen. Only nonspecific staining of chondrocytes was seen on untreated BNC sections stained with antibodies to 1α 2α 3α collagen. However, after treatment with hyaluronidase, diffuse staining of the territorial matrix was obtained (Fig. 11). This was considered to be specific, as the staining was abolished by blocking controls.

C-PS 1 and C-PS 2 collagens. The staining patterns for these 2 collagens were similar to that for 1α 2α 3α collagen.

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**Fig. 7** Control. Untreated BNC section overlaid with normal rat serum (1/10 dilution), washed, treated with FITC-labelled rabbit anti-rat immunoglobulins, and washed again before mounting. (×470).

**Fig. 8** Control. Untreated BNC section overlaid with rat anti-type II collagen, washed, overlaid with unconjugated rabbit anti-rat immunoglobulins (1/5 dilution), washed and overlaid with FITC-labelled rabbit anti-rat immunoglobulins, and washed again before mounting. (×470).

**Fig. 9** BNC treated with hyaluronidase. Immunolocalisation of type II collagen. (×470).

**Fig. 10** BNC treated with bacterial collagenase, and stained for type II collagen. (×470).
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collagen (Figs 12 and 13). Staining was abolished by blocking controls.

*Type V collagen.* Again, only nonspecific staining of chondrocytes was seen on untreated BNC sections. On hyaluronidase-treated sections very bright, discrete pericellular staining was observed (Fig. 14). This was considered specific by the criteria of using normal guinea-pig serum and blocking controls.

*Types I, III, and IV collagen.* No staining was seen with antibodies to these 3 collagens, either on untreated BNC or on sections treated with hyaluronidase.

*Fibronectin.* Anti-fibronectin antibodies stained the interterritorial matrix on both untreated and hyaluronidase-treated sections (Fig. 15), and this was shown to be specific by controls with normal rabbit serum and blocking controls. No staining of the pericellular and territorial region was seen.

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**Fig. 11** BNC treated with hyaluronidase. Immunolocalisation of 1α 2α 3α collagen. (×470).

**Fig. 12** BNC treated with hyaluronidase. Immunolocalisation of C-PS 1 collagen. (×470).

**Fig. 13** BNC treated with hyaluronidase. Immunolocalisation of C-PS 2 collagen. (×470).

**Fig. 14** BNC treated with hyaluronidase. Immunolocalisation of type V collagen. (×470).

**Fig. 15** BNC, untreated. Immunolocalisation of fibronectin. (×470).
This study suggests that different collagen types are distributed in defined areas within the cartilage matrix. Type II collagen, the most abundant collagen type in cartilage, appears on untreated sections to be distributed interterritorially. However, after treatment with hyaluronidase it can be seen that type II collagen is present throughout the cartilage matrix. In contrast, 1α 2α 3α collagen and the C-PS collagens cannot be detected on untreated sections of BNC and can only be visualised after enzyme treatment to remove proteoglycans. It would appear that these collagens are located around the chondrocytes, in the territorial zone, and that they are either absent, or present in undetectable quantities, in the interterritorial spaces.

Type V collagen presents an interesting anomaly. Although there is one report of type V collagen being biochemically detected in cartilage, this finding has not been confirmed by us using biochemical methods. The type V antiserum was extensively absorbed against other collagens and gave the expected distribution on placental and other tissues (Evans and Abedin, unpublished observations), so it seems unlikely that the poor specificity of this reagent was responsible for an artefactual result. Immunofluorescent studies are renowned for the production of 'edge' effects, where nonspecific staining appears on the edge of sections. It could be argued that such an effect was observed with the type V antiserum, but this is unlikely, as the immunofluorescence was abolished by blocking controls. Two possibilities could explain the staining with the type V antiserum; the antibodies may cross-react with a hitherto undiscovered collagen present in cartilage which is present in low quantities or is intimately associated with other constituents of cartilage. Alternatively, type V collagen may indeed be present in cartilage, and its detection by immunohistochemical means but not by biochemical methods may reflect the differing sensitivities of the 2 techniques. It is interesting to note that the fluorescent patterns on cartilage which have been described with sera from patients suffering from relapsing polychondritis are similar to that observed with our type V antiserum. Previously these antibodies have been assumed to be directed against type II collagen.

Antibodies to fibronectin failed to stain frozen sections of the cartilage matrix of chick sternal cartilage. However, these authors showed that chondrocytes from this cartilage in culture synthesise and secrete fibronectin, indicating that these cells do have the potential to produce fibronectin under certain conditions. Our study suggests that fibronectin is present in adult bovine nasal cartilage matrix. It locates only in the interterritorial matrix, in the areas from which 1α 2α 3α and the C-PS collagens appear to be absent. The only type of collagen we have detected in this region is type II.

Poole et al. have shown in an immunohistochemical study of proteoglycans and link protein of bovine articular cartilage that these have well-defined locations. Proteoglycan was evenly distributed throughout the cartilage matrix, yet more link protein was detectable in the interterritorial sites in the middle and deep zones of the bovine articular cartilage. Well-defined zones of weaker territorial staining for link protein stained strongest for chondroitin sulphate.

Our study shows that collagen types have well-defined locations in bovine nasal cartilage. The functional significance of this is not yet clear. C-PS 1 and C-PS 2 collagens probably exist in a much larger form in the intact tissue. Moreover, they could be specifically associated with the 1α 2α 3α collagen, since all 3 collagens have a similar location in the territorial zone. Such an association could provide the protective enclosures believed by several workers to exist around the chondrocytes. An extension of this work at the level of the electron microscope might be more informative with regard to this suggestion, which at present is purely speculative.

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References

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