Immunochemical and biochemical comparisons between embryonic chick bone marrow and epiphyseal cartilage chondroitin/dermatan sulphate proteoglycans

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Summary

Chondroitin sulphate proteoglycans obtained from embryonic chick bone marrow and epiphyseal cartilage were compared using immunochemical and biochemical analyses. Proteoglycans from each tissue, separated on CsCl density gradients, under dissociative conditions, into high (1·6 g ml⁻¹), medium (1·5 g ml⁻¹) and low (1·4 g ml⁻¹) buoyant density fractions, were immunochemically analysed, using a panel of monoclonal antibodies that specifically recognize chondroitin 4-/dermatan sulphates, chondroitin 6-sulphate, keratan sulphate, the hyaluronate binding region present on connective tissue proteoglycans, and link protein. The same antibodies were used in Western blot analyses to detect intact proteoglycan monomers and core proteins that had been fractionated by agarose–polyacrylamide and by sodium dodecyl sulphate–polyacrylamide gel electrophoresis. Specific differences between marrow and cartilage proteoglycans were detected. In CsCl gradients, marrow proteoglycans displayed a higher degree of heterogeneity in terms of buoyant densities and hexuronate distribution. Keratan sulphate chains were constituents of the majority of 'large' proteoglycans in the marrow; however, a portion of the large proteoglycans in marrow middle buoyant density fraction either lacked keratan sulphate chains or were substituted with a form different from that found on cartilage proteoglycans. Marrow lacked 'small' chondroitin/dermatan sulphate proteoglycans that were present in cartilage and contained a more heterogeneous population of proteoglycans, particularly in the lower buoyant density fractions. Both marrow and cartilage were similar in that they contained, as their major components, large, aggregating proteoglycans and link proteins that were immunochemically and biochemically identical. The significance of these differences between marrow and cartilage proteoglycans remains to be determined, but they may, in part, be responsible for imparting unique characteristics to the haematopoietic extracellular matrices.

Key words: proteoglycans, marrow, cartilage.

Introduction

The concept of a haematopoietic inductive microenvironment was first postulated by Trentin and his colleagues (Curry & Trentin, 1967; Curry et al. 1967; Trentin, 1978). Since then, it has become increasingly evident that non-haematopoietic stromal cells are largely responsible for generating this environment (Dexter, 1982). Stromal cells appear, in large part, to be responsible for elaborating an extracellular matrix milieu that may be critical for haematopoietic–stromal cell interactions (Spooner et al. 1983; Zuckerman & Wicha, 1983; Zipori et al. 1985).

A diverse group of molecules, proteoglycans (Hascall & Hascall, 1981; Hassell et al. 1986), are a major component of hematopoietic extracellular matrices. The amounts and relative compositions of these proteoglycans have been shown to vary according to haematopoietic status (Noordegraaf & Ploemacher, 1979; Noordegraaf & Ploemacher, 1980; Noordegraaf et al. 1981). Gallagher et al. (1983) provided clear evidence for differences between proteoglycans associated with stromal cells and proteoglycans released into the medium in long-term haematopoietic cultures. Cell-associated proteoglycans were enriched in heparan sulphate and also contained some chondroitin sulphate, whereas those released into the medium consisted almost entirely of chondroitin sulphates. This work was confirmed and extended by Wight et al. (1986), who have also provided important information concerning the nature of intact proteoglycan molecules in hematopoietic cultures. All previous
studies were limited to the nature of the glycosaminoglycan chains.

The importance of studying intact proteoglycan molecules has been emphasized by recent studies that implicate tissue-specific proteoglycans with the ability to sequester growth factors selectively (Okai, 1985; Gordon et al. 1987; Vlodavsky et al. 1987). Sequestration and concentration of growth factors on surfaces of stromal cells would account, in part, for the difficulties in recovering haematopoietic growth factors from long-term marrow cultures (Dexter et al. 1977; Williams et al. 1977) and might, in part, account for the apparent requirement for close cell-to-cell contact between haematopoietic and stromal cells (Dexter, 1982; Bentley, 1981).

In the present study, embryonic chick marrow chondroitin sulphate proteoglycans were characterized and compared with embryonic chick cartilage proteoglycans, about which considerable information has accumulated (Kimata et al. 1974; Vasan & Lash, 1978; Noro et al. 1983; Shinomura et al. 1983; Carrino & Caplan, 1985). Well-characterized monoclonal antibodies that specifically recognize chondroitin 4-sulphate and dermatan sulphate (2B6), chondroitin 6-sulphate (3B3), and keratan sulphate (5D4) chains were used to identify GAG chains and to immunolocate proteoglycans on Western blots after composite agarose-polyacrylamide gel electrophoresis of intact monomers and sodium dodecyl sulphate-polyacrylamide gel electrophoresis (SDS-PAGE) analyses of core proteins (Caterson et al. 1983, 1985, 1986, 1987). Monoclonal antibody 1C6, which specifically recognizes the hyaluronate binding region, was used to identify chondroitin sulphate proteoglycans that have the potential to form aggregates (Stevens et al. 1984; Caterson et al. 1986, 1987), while antibody 8A4 was used to identify link protein (Caterson et al. 1986), also a component of macromolecular aggregates (Hay, 1981). These methods provided a highly sensitive means of examining the total proteoglycan composition of these tissues, not merely the newly synthesized pool obtained after metabolic labelling with radioactive tracers.

A heterogeneous population of marrow chondroitin sulphate proteoglycans was identified, many of which were nearly identical to cartilage proteoglycans. However, comparative studies indicated specific differences in two of the three proteoglycan fractions. These results target chondroitin sulphate proteoglycan species that may be marrow-specific.

### Materials and methods

#### Antibodies

All monoclonal antibodies used in these studies had specificities directed against different components of chondroitin sulphate proteoglycans. See Table 1 for details concerning epitopes recognized by these monoclonal antibodies.

#### Extraction and buoyant density fractionation of proteoglycans

Femora and tibiae were dissected from cold anaesthetized day 17 chick embryos (75 embryos per experiment) and stripped of soft tissue. The epiphysial cartilages were removed and the bones cut lengthwise in half. Marrow was scraped from the bones directly into cold 4 M-guanidine·HCl containing protease inhibitors (Sajdera & Hascall, 1969). After all marrow had been removed, excess guanidine·HCl was added so that the final volume was at least 20 times that of marrow. This was stirred continuously for 24 h at 4°C, centrifuged at 15 000 g for 30 min, and the supernatants were collected. Solid CsCl was added to give a density of 1.5 g ml⁻¹. After centrifugation at 100 000 g at 10°C for 48 h under 'dissociative' conditions (Oegema et al. 1975), the bottom (MD1 or CD1), middle (MD2 or CD2) and top (MD3 or CD3) thirds of the CaCl₂ density gradients from the marrow or cartilage extracts, respectively, were carefully collected and small samples were taken for density and hexuronic acid analysis (Carney et al. 1986). Each fraction was dialysed exhaustively against 0.15 M-NaCl and then distilled water. Dry weights were determined and samples stored as lyophilized powder.

#### Dot-blot analysis

Marrow and cartilage proteoglycans from each fraction were dissolved in 4 M-guanidine·HCl, 0.001 M-EDTA, 0.1 M-Tris-acetate, pH 8.0, to give a concentration of 1 mg ml⁻¹. Samples were divided in half: one part was treated with 0.065 M-dithiothreitol for 4 h at 50°C (reduced), the other part was incubated at 50°C without dithiothreitol (native). Both portions were treated with 0.16 M-isoadic acid in the dark at room temperature for 20 min then dialysed against three changes of phosphate-buffered saline containing azide (PBS-AZ) over a 24-h period (Stevens et al. 1984).

Native and reduced and alkylated samples were applied by vacuum suction to same sheet of nitrocellulose using a Schleicher and Schuell Minifold apparatus (Caterson et al. 1987). All nitrocellulose sheets were incubated in 5% bovine serum albumin (BSA) at room temperature for 1 h and then incubated with 0.01 unit ml⁻¹ of chondroitinase ABC prior to immunoloccation. Each monoclonal antibody was diluted 1/100 in 5% BSA in 0.1 M-Tris–saline and was incubated with antigen on nitrocellulose sheets for 90 min at room temperature. The first antibody incubation step was eliminated for control

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**Table 1. Monoclonal antibodies used for this study and their epitopes**

<table>
<thead>
<tr>
<th>Antibody</th>
<th>Epitope</th>
<th>Reference</th>
</tr>
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<tbody>
<tr>
<td>2B6</td>
<td>Delta-unsaturated 4-sulphated epitope generated by chondroitinase ABC digestion of chondroitin 4-/dermatan sulphates</td>
<td>Caterson et al. (1985)</td>
</tr>
<tr>
<td>3B3</td>
<td>Delta-unsaturated 6-sulphated epitope generated by chondroitinase ABC digestion of chondroitin 6-sulphate</td>
<td>Caterson et al. (1985)</td>
</tr>
<tr>
<td>5D4</td>
<td>Linear, sulphated heptasaccharide, or larger, of poly-N-acetyllactosamine (keratan sulphate)</td>
<td>Mehnert et al. (1986); Caterson et al. (1983)</td>
</tr>
<tr>
<td>1C6</td>
<td>Hyaluronate binding region of core proteins after reduction and alkylation</td>
<td>Caterson et al. (1986)</td>
</tr>
<tr>
<td>8A4</td>
<td>Link protein</td>
<td>Caterson et al. (1986)</td>
</tr>
</tbody>
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samples. After washing three times with Tris-saline, peroxidase-conjugated goat anti-mouse second antibody diluted 1/500 was incubated in each sheet. After washing with Tris-saline, 4-chloro-l-naphthol and H₂O₂ substrate was added and coloured precipitating products developed. The reaction was stopped by washing with PBS-AZ. The nitrocellulose sheets were air-dried prior to photographic reproduction.

**Fractionation of intact proteoglycan monomers on composite agarose-polyacrylamide gels**

Composite agarose-polyacrylamide gels were run using a modified method described by Carney et al. (1986). Agarose (0-6 % final concentration) was suspended in gel buffer (0-04 M-Tris-acetate/0-001 M-Na₂SO₄, pH 6.8) and heated to 90°C before adding acrylamide/bisacrylamide (1-2 % final concentration). The final mixture was poured into preheated (50°C) slab gel plates and allowed to set at 4°C for 1 h. Plates were equilibrated overnight in 4 M-urea and gel buffer (Carney et al. 1986). Samples dissolved in 8 M-urea were applied to the gel and electrophoresed at constant voltage for 2 h until the tracking dye had migrated 3-4 cm. Some gels were stained with 0-02 % Toluidine Blue to reveal proteoglycans. Samples from other gels were immunolocated after transfer onto nitrocellulose (see dot-blot procedures). Enzyme digestion, when needed, was performed on nitrocellulose-bound samples prior to immunolocation. The first antibody incubation step was omitted for one set of samples. Antibodies as a negative control, the first antibody incubation step was eliminated for one set of samples.

Western blot analysis of link protein was performed as described above, except that antibody 8A4 was used for immunolocation after 10 % SDS-PAGE analyses of the MD3 and CD3 fractions.

**Results**

**Comparison of marrow and cartilage proteoglycans fractionated on dissociative CsCl density gradients**

Proteoglycan monomers, extracted from the same day 17 embryonic chick femoral and tibial bone marrows and epiphyseal cartilages, were compared using combined immunochemical and biochemical analyses. Proteoglycan monomers from both tissues were fractionated on CsCl density gradients under dissociative conditions and three equal portions were collected (Table 2). Marrow proteoglycans differed from cartilage proteoglycans in that a substantially lower proportion of the total hexuronate (56 % for MD1 versus 75 % for CD1) was found in the high buoyant density (D1) fractions and correspondingly higher proportions of hexuronate were found in the lower buoyant density fractions, MD2 and MD3. Marrow contained a substantially higher proportion of extractable material that partitioned to the low buoyant density fraction (68 % dry weight compared to 28 % dry weight, Table 2), indicating that marrow contained a higher content of non-proteoglycan molecules than did cartilage. Both D1 fractions were identical in terms of their hexuronate to dry weight ratio and buoyant densities, but the hexuronate to dry weight ratio of marrow fraction MD2 was lower than that of marrow fraction MD1, as well as cartilage fractions CD1 and CD2. This implies that MD2 proteoglycans were probably substituted with either fewer or shorter GAG chains than those of proteoglycans in all other D1 and D2 fractions. The presence of partially degraded proteoglycans in marrow might also account for this difference.

**Immunochemical comparisons of marrow and cartilage proteoglycan monomers**

Epitopes recognized by three carbohydrate-specific monoclonal antibodies (Table 1) were found in each marrow and cartilage fraction (Fig. 1). This suggests that...
Fig. 1. Dot-blot analysis was used to compare marrow MD3 (1), MD2 (2), MD1 (3), CD3 (4), CD2 (5), and CD1 (6) fractions. Samples from each fraction were immunolocated with: A, monoclonal antibody 2B6, specific for chondroitin 4-/dermatan sulphates; B, antibody 3B3, specific for chondroitin 6-sulphate, antibody 5D4; C, specific for keratan sulphate; D, antibody 1C6, specific for epitopes associated with hyaluronate binding region, and, as a control; E, no first antibody. Antigen in the top rows of each panel were unreduced, while proteoglycans in the bottom rows were reduced and alkylated prior to immunoanalyses.

Fig. 2. Samples from low buoyant density marrow and cartilage fractions were analysed by SDS-PAGE, under reducing conditions, on a 10% gel, electroblotted onto nitrocellulose, and immunolocated using antibody 8A4, specific for link protein. One major band, representing a protein with an apparent molecular weight of \(43 \times 10^3\), was immunolocated in both marrow (lane a) and cartilage (lane b) fractions. Also, a second band, probably representing a second and minor link protein, was weakly immunolocated (thin arrows) in each fraction. Marrow (lane a) contained a faster migrating peptide (thick arrow), which was also immunolocated by antibody 8A4.

Low buoyant density fractions from both marrow and cartilage contained one major link protein, as identified by link protein-specific antibody 8A4 (Fig. 2). This link protein migrated on a 10% SDS–polyacrylamide gel with an apparent molecular weight of \(43 \times 10^3\). In addition, a minor, slightly faster migrating link protein was present in both tissues. The narrow fraction contained a faster migrating peptide that was also immunolocated by antibody 8A4. These studies were repeated using several different marrow and cartilage samples, comparing the results both with and without mercaptoethanol reduction. The faster migrating peptide was never identified in non-reduced samples, but was identified in some
Fig. 3. Serial dilutions of intact proteoglycans from marrow and cartilage fractions MD1 (lanes a), CD1 (lanes b) and, as a standard, canine femoral articular cartilage proteoglycan monomer (lanes c) were analysed by electrophoresis on composite agarose–polyacrylamide gels. The gel was stained with 0.2% Toluidine Blue.

of the reduced samples. Previously, it had been shown that reduction by mercaptoethanol was needed to identify link protein fragments (Mort et al. 1985). Thus, it is possible that the small peptide is a proteoglycally cleaved component of intact link protein. Identification of both link protein and hyaluronate binding region is consistent with the presence of aggregating proteoglycans in bone marrow, which are similar to those found in cartilage (Hascall & Hascall, 1981).

Comparison of intact chondroitin sulphate proteoglycan monomers from marrow and cartilage using composite agarose–polyacrylamide gels

Marrow and cartilage high buoyant density proteoglycans were analysed according to their electrophoretic mobilities in composite agarose–polyacrylamide gels. Using this procedure, the heterogeneous proteoglycan subpopulations can be separated and then analysed biochemically using immunochemical procedures (Carney et al. 1986). In Fig. 3, marrow and cartilage proteoglycans, along with a canine cartilage proteoglycan monomer standard, were identified by staining the gel with Toluidine Blue. Proteoglycans from both chick tissues migrated with similar mobilities that were slower than the canine standard. A slower migration in composite agarose–polyacrylamide gels is usually indicative of large chondroitin sulphate-containing proteoglycans with relatively low amounts of keratan sulphate. In both tissues the proteoglycan bands were diffuse, consistent with a high degree of polydispersity in size and charge.

Electrophoretic analysis on composite agarose–polyacrylamide gels was repeated, but now proteoglycans were identified on Western blots using monoclonal antibodies 2B6 and 3B3 separately (Fig. 4). Chondroitin 6-sulphate-specific antibody 3B3 identified a slower migrating subpopulation (Fig. 4, lanes a and b). In contrast, antibody 2B6, specific for chondroitin 4-/dermatan sulphates detected a faster migrating subpopulation of proteoglycans in both marrow and cartilage (Fig. 4, lanes a' and b'). These data suggest that at least two subpopu-
Fig. 5. Proteoglycan core protein derived from chondroitinase ABC-digested marrow and cartilage fractions MD1 (lane a), CD1 (lane b), MD2 (lane c), CD2 (lane d), MD3 (lane e), and CD3 (lane f) were analysed by SDS–PAGE on a 5 % gel, electroblotted onto nitrocellulose, and immunolocated with a mixture of monoclonal antibodies 2B6 and 3B3. Molecular weight markers for 212 and 97(×10^3)M_r are indicated. Arrowheads in lane e indicate three pairs of bands immunodetected in the MD3 fraction, while the arrowhead in lane f indicates the single band immunodetected in the CD3 fraction.

since these molecules were not effectively resolved in the running gel. Small proteoglycan core proteins with apparent molecular weights of 43 and 47(×10^3) were also immunochemically identified in cartilage D2 fractions after electrophoresis on a 7 % to 17 % linear gradient SDS–polyacrylamide gel, but similar core proteins were not detected in the equivalent marrow fraction (Fig. 6A,B), thus indicating a difference in the proteoglycan compositions of these two tissues.

Intermediate-sized core proteins, migrating with apparent molecular weights ranging from 100 to 200(×10^3), were detected on a Western blot using a mixture of antibodies 2B6 and 3B3 (Fig. 5, lanes e and f). Multiple bands were detected in the marrow fraction MD3 (lane e). In contrast, only one such band, with an apparent molecular weight of 120×10^3 and corresponding to the second fastest migrating MD3 band, was identified in the cartilage fraction CD3 (lane f). It is likely that some or possibly all of the proteoglycan core proteins identified in the low buoyant density fractions represented partial degradation products of large proteoglycans.

Comparative dot-blot immunoanalyses of marrow and cartilage fractions (Fig. 1) revealed the presence of keratan sulphate epitopes, using antibody 5D4, in both marrow and cartilage proteoglycans. As noted above, the MD2 fraction contained relatively less of this epitope than did any other marrow or cartilage fractions. The low level of keratan sulphate substitution of the MD2 fraction was further confirmed on Western blots (Fig. 7). Anti-

body 5D4 immunolocated epitopes in both marrow (lane a) and cartilage (lane b) high buoyant density fractions, but failed to identify epitopes in the marrow middle density fraction (lane c), although epitopes were identified in the CD2 fraction (lane d). The discrepancy

Fig. 6. Proteoglycan core protein derived from chondroitinase ABC-digested marrow and cartilage fractions MD2 (lane a) and CD2 (lane b) were analysed by SDS–PAGE on a 7 % to 17 % linear gradient gel, electroblotted onto nitrocellulose, and immunolocalized with a mixture of monoclonal antibodies 2B6 and 3B3. Arrowheads indicate 43 and 47(×10^3)M_r bands in CD2 fractions.

Fig. 7. Proteoglycan core protein derived from chondroitinase ABC-digested marrow and cartilage fractions MD1 (lane a), CD1 (lane b), MD2 (lane c), and CD2 (lane d) were analysed by SDS–PAGE on a 5 % gel, electroblotted onto nitrocellulose, and immunolocalized with monoclonal antibody 5D4, specific for keratan sulphate. The 212×10^3 M_r marker is indicated.
between these results and those shown in Fig. 1 are probably related to the lower sensitivity of Western blot analyses compared with that of dot-blot analyses. Repetition of these studies using the same and different keratan sulphate-specific antibodies produced the same results as those indicated above. These results indicate that some of the marrow proteoglycans differ, in terms of their constituent glycosaminoglycan chains, from cartilage proteoglycans.

Discussion

The present data indicate that, while there was a high degree of similarity between chick marrow and cartilage proteoglycans, diversity also existed. Several specific differences between marrow and cartilage proteoglycans were noted. (1) In marrow, proteoglycan monomers appeared to display a higher degree of heterogeneity. A substantially higher proportion of proteoglycans resided in the middle and low buoyant density fractions. (2) Marrow middle buoyant density proteoglycans were substituted with either less or a different form of keratan sulphate. (3) Small proteoglycans, characteristic of cartilage, bone, and some other non-cartilaginous tissues (Shinomura et al. 1983; Hassell et al. 1986), appeared to be absent from marrow. (4) Finally, marrow contained what appeared to be a heterogeneous population of proteoglycans in the low buoyant density fraction that were not detected in the equivalent cartilage fraction.

Marrow from chick embryos contained large chondroitin sulphate proteoglycan monomers, which possessed large core proteins similar in size to those found in cartilage. Nonetheless, marrow proteoglycans apportioned differently from cartilage proteoglycans on dissociative CsCl gradients. Marrow contained a substantially higher proportion of intermediate and low buoyant density proteoglycans. This suggested that differences in glycosylation and/or sulphation patterns might have existed between some of the marrow and cartilage proteoglycans. Partial degradation of proteoglycans would also result in smaller sizes and lower buoyant densities. Hence, differences in endogenous turnover of proteoglycan could affect the partitioning of proteoglycans on CsCl gradients.

A high proportion of marrow proteoglycans were cartilage-like in terms of their large size, presence of an epitope associated with the hyaluronate binding region, and substitution with keratan sulphate. Nevertheless, some of the marrow proteoglycans, particularly those in the MD2 fraction, differed from cartilage proteoglycans in regard to keratan sulphate substitution. Antibody 5D4 provided a weak immunolocalization of MD2 when the more sensitive dot-blot assay was used, and failed to identify epitopes on MD2 on less-sensitive Western blots after SDS–PAGE fractionation. These data suggest that relatively little keratan sulphate was associated with marrow MD2 proteoglycans. Alternatively, they suggest that keratan sulphate, if present, did not contain epitopes that were readily recognized by antibody 5D4. Chemical desulphation of keratan sulphate chains reduced antigenicity for this antibody (Mehmet et al. 1986; Funderburgh et al. 1987). Thus, low sulphated keratan sulphate might be present on some marrow proteoglycans, particularly those in the MD2 fraction, and this might account for the weak immunochromic reactivity of antibody 5D4. Marrow also differed from cartilage in that small chondroitin sulphate proteoglycans were absent, or at least undetected, by the methods used in this study. Small proteoglycans, with core proteins of 43 and 47×10^3 kDa, as assessed by SDS–PAGE, have been identified in cartilages obtained from a variety of vertebrate species, including those from embryonic chicks (Shinomura et al. 1983; Hassell et al. 1986). It was unlikely that this was due to procedural problems, since two core proteins with apparent molecular weights of 43 and 47×10^3 were detected in the CD2 fraction. Small proteoglycans, usually substituted with dermatan sulphate chains, are not entirely restricted to cartilages (Hassell et al. 1986). Wight et al. (1986) identified, as a minor species, a small dermatan sulphate proteoglycan produced by long-term cultures of human marrow. These ‘small’ proteoglycans, in a variety of tissues, often appear to be closely associated with collagen types I and III (Hassell et al. 1986). The relative sparsity of such collagens in embryonic chick marrow might account for the absence of small proteoglycans in chick marrow (Sorrell et al. 1987).

Marrow, but not cartilage, contained an apparently heterogeneous population of proteoglycans in the low buoyant density fraction. At least six core proteins were detected on Western blots of marrow D3 proteoglycans that had been fractionated by SDS–PAGE. The apparent molecular weights of marrow core proteins ranged from 100 to 200×10^3 kDa in marrow, compared to only one core protein, of about 120×10^3 kDa, in cartilage. These putative, intermediate-sized proteoglycans have yet to be identified definitively. At present two possibilities exist: one or more bands in the MD3 fraction might represent the core protein(s) of a uniquely different proteoglycan; on the other hand, these might be partially degradation products of large core protein(s), those larger than 212×10^3 kDa. Both marrow and cartilage proteoglycans were extracted simultaneously in the same media containing protease inhibitors. Multiple bands were absent from the equivalent cartilage fraction (a tentative identification can be made for the one band in the CD3 fraction, see below). Thus, it is unlikely that degradation, if it existed, could be attributed to the extraction procedures. Rather, if degradation occurred, it is more likely to be a consequence of higher endogenous turnover of marrow proteoglycans.

Proteoglycans comprise a diverse group of molecules (Hassell et al. 1986). Nevertheless, this diversity is limited. Therefore, it is not surprising that similarities among marrow and cartilage proteoglycans exist. Wight et al. (1986) also commented on the overall similarity of proteoglycans obtained from long-term human marrow cultures to proteoglycans obtained from aorta, and in this study, similarities among chick marrow and cartilage proteoglycans were clearly evident. As with cartilage, most marrow proteoglycans belong to a family of large...
proteoglycans (Heinegard et al. 1985; Hassell et al. 1986), some of which, particularly those in the high buoyant density fraction, contained a hyaluronate binding region. This implies that these proteoglycans are capable of associating with hyaluronic acid to form macromolecular aggregates (Hascall & Hascall, 1981). Further evidence for the presence of proteoglycan aggregates was provided by the immunodetection of link protein, also known to be a component of macromolecular aggregates (Hay, 1981). Similar proteoglycans have been identified in a wide variety of non-cartilaginous tissues and cultured fibroblastic cell lines (Nerling et al. 1978; McMurtrey et al. 1979; Wagner et al. 1983; Schafer et al. 1984; Heinegard et al. 1985; Hassell et al. 1986). Indirect evidence for proteoglycan aggregates was obtained from histochemical studies of chick marrow where the high iron diamine–silver proteinate method was used to stain chondroitinase-ABC-digestible material and hyaluronidase, which is known to degrade hyaluronate specifically (Hay, 1981), resulted in the loss of sulphated glycoconjugates, presumably chondroitin sulphates bound to hyaluronate. The presence of hyaluronate in marrow and other haematopoietic tissues is well documented (Olsson, 1971; Taniguchi et al. 1974; Noordegraaf & Ploemacher, 1979, 1980; Noordegraaf et al. 1981; Wight et al. 1986; Gordon et al. 1987; Óguri et al. 1987). This does not mean that all marrow proteoglycans participate in macromolecular aggregates. On the basis of inferences obtained from dot-blot analyses, the middle and low buoyant density marrow proteoglycans did not contain a hyaluronate binding region and consequently were unlikely to be capable of forming aggregates. If so, this would place the large core protein of the proteoglycans of the middle buoyant density marrow and cartilage fractions in the family of large non-aggregating proteoglycans (Heinegard et al. 1985).

The occurrence of keratan sulphate proteoglycans in marrow was unexpected, since chondroitin sulphate proteoglycans containing keratan sulphate chains have been considered exclusive to cartilage. Keratan sulphate in the extracellular spics of chick marrow also has been identified using immunocytochemical methods at the electron-microscopic level (Sorrell et al. 1988). Recently, keratan sulphate chains have been identified in tissues other than cartilage and cornea. Circumstantial evidence for its presence has been presented for aorta (Heinegard & Hascall, 1979), and more recent immunocytochemical evidence has been presented for its presence in developing conjunctiva (Sundarraj et al. 1986) and in the corpora amylacea of the human brain (Liu et al. 1987). Funderburgh et al. (1987) have surveyed a variety of bovine and chick non-cartilaginous tissues and have found that keratan sulphate substitution is much more widespread than anticipated. The function of keratan sulphate-substituted proteoglycans in non-cartilaginous tissues, including marrow, remains to be established.

Only one core protein, with an apparent molecular weight of 120X10^2, was identified in the cartilage low buoyant density fraction. This proteoglycan has yet to be positively identified, but it is possible that it represents PG-Lt described by Noro et al. (1983), which has subsequently been renamed collagen type IX (Hassell et al. 1986; Irwin & Mayne, 1986). Huber et al. (1986) have estimated the size of this, minus the chondroitin sulphate chain, at 115X10^2. If the 120X10^2 band proves to be collagen type IX, then it is also probably present in marrow.

The demonstration that there were apparent differences in the composition of chondroitin sulphate proteoglycan between marrow and cartilage may prove to be significant. Chondroitin sulphates have been demonstrated to interact with some types of blood cells, particularly lymphocytes, with a consequent modulation of some of their cellular functions (Darzynkiewicz & Balazs, 1971; Siegel et al. 1987). Extracellular matrix and/or components associated with matrix have been shown to promote haematopoiesis either directly or indirectly. Matrix derived from haematopoietic tissues, but not other tissues, promotes the attachment of haematopoietic stromal cells to culture dishes and consequently promotes growth and differentiation of haematopoietic cells (Campbell et al. 1985). Similarly, bone marrow matrix has been shown to alter the phenotypic expression of HL-60 cells (Luijktart et al. 1987). It is possible that these functions are, at least in part, mediated by the apparent ability of extracellular matrix components, particularly proteoglycans, to sequester haematopoietic growth factors (Okai, 1985; Gordon et al. 1987). The implication to be drawn from these studies is that haematopoietic-specific matrix components, probably proteoglycans, may be involved in regulating haematopoiesis. If so, it should be possible to identify those proteoglycans. The present study provides evidence that some marrow chondroitin sulphate proteoglycans differ from those found in cartilage. Whether these proteoglycans in marrow have functional roles in haematopoiesis remains to be determined.

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References


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