Abstract. The glycosaminoglycan chains of cell surface heparan sulfate proteoglycans are believed to regulate cell adhesion, proliferation, and extracellular matrix assembly, through their interactions with heparin-binding proteins (for review see Ruoslahti, E. 1988. Annu. Rev. Cell Biol. 4:229–255; and Bernfield, M., R. Kokenyesi, M. Kato, M. T. Hinkes, J. Spring, R. L. Gallo, and E. J. Lose. 1992. Annu. Rev. Cell Biol. 8:365–393). Heparin-binding sites on many extracellular matrix proteins have been described; however, the heparin-binding site on type I collagen, a ubiquitous heparin-binding protein of the extracellular matrix, remains undescribed. Here we used heparin, a structural and functional analogue of heparan sulfate, as a probe to study the nature of the heparan sulfate proteoglycan-binding site on type I collagen. We used affinity coelectrophoresis to study the binding of heparin to various forms of type I collagen, and electron microscopy to visualize the site(s) of interaction of heparin with type I collagen monomers and fibrils. Using affinity coelectrophoresis it was found that heparin has similar affinities for both procollagen and collagen fibrils ($K_d$'s $\sim$60–80 nM), suggesting that functionally similar heparin-binding sites exist in type I collagen independent of its aggregation state. Complexes of heparin–albumin–gold particles and procollagen were visualized by rotary shadowing and electron microscopy, and a preferred site of heparin binding was observed near the NH$_2$ terminus of procollagen. Native or reconstituted type I collagen fibrils showed one region of significant heparin–gold binding within each 67-nm period, present near the division between the overlap and gap zones, within the “a” bands region. According to an accepted model of collagen fibril structure, our data are consistent with the presence of a single preferred heparin-binding site near the NH$_2$ terminus of the collagen monomer. Correlating these data with known type I collagen sequences, we suggest that the heparin-binding site in type I collagen may consist of a highly basic triple helical domain, including several amino acids known sometimes to function as disaccharide acceptor sites. We propose that the heparin-binding site of type I collagen may play a key role in cell adhesion and migration within connective tissues, or in the cell-directed assembly or restructuring of the collagenous extracellular matrix.

Interactions between heparan sulfate proteoglycans (HSPGs) and collagen fibrils are thought to play important roles in cell adhesion (Koda et al., 1985; Piepkorn and Chapman, 1985), and in the formation of extracellular matrix structures such as basement membranes (Tillibary et al., 1988, 1990; Koliakos et al., 1989). It has been shown that HSPGs, heparan sulfates, and heparin each bind to type I collagen fibrils (Stamataglou and Keller, 1982; Koda et al., 1985; Keller et al., 1986; LeBaron et al., 1989).

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necessary for HSPG or heparin binding remain unknown. Thus, here we attempted to define the HSPG-binding sites on type I collagen, owing to the likely importance of these sites in the interactions between cell surfaces and type I collagen—one of the most abundant molecules of the extracellular matrix. Thus, in this manuscript we have characterized heparin–collagen interactions by affinity coelectrophoresis (ACE) and, using electron microscopy, sites of interaction were mapped for the binding of heparin–collodial gold complexes to individual type I procollagen molecules and to both native and reconstituted type I collagen fibrils.

Materials and Methods

Collagen Preparations

Procollagen Type I. Procollagen type I was isolated as detailed (Olsen et al., 1976). Briefly, leg tendons were dissected from 17-d-old chick embryos, digested with trypsin and collagenase, and then the tendon fibroblasts were pelleted. Tendons from twelve dozen embryos yielded about 2 × 10^9 cells, which were placed in suspension culture for 6 h. Procollagen type I was isolated from the culture medium of the suspension cultures by ammonium sulfate precipitation with final protein yields of ∼1.5 mg observed. Procollagen type I purity was confirmed by SDS-PAGE analysis, and protein concentration was determined by amino acid analysis, performed by the Biochemistry Laboratory of the Brigham and Women's Hospital (Boston, MA).

Collagen Type I

Acid Soluble. Collagen from rat tail tendons was isolated as described (San Antonio et al., 1992). Briefly, tendons were subjected to three cycles of extraction/precipitation with 0.5 N acetic acid/7.5% NaCl, and purity was confirmed by SDS-PAGE.

Native. Tendons were removed from the tails of 5–10 Long Evans rats and were placed in 0.1 M Tris HCl, 0.1 M NaCl, pH 7.5, with 1 mM PMSF and 1 mM EDTA. All procedures were conducted at 4°C, and used the same buffer as detailed above. Tendons were rinsed several times in buffer, and were diced into pieces of ∼1 cm using a razor blade and scissors. The sample was suspended in 100 ml of buffer, and homogenized at high speed in homogenization buffer, and then used in electron microscopy studies. Some samples were further treated with chondroitinase ABC. Fibril suspensions were diluted through these gels was as described above.

Preparation of Samples for Electron Microscopy of Procollagen–Heparin–Gold Complexes

Procollagen was dialyzed into 0.2 M potassium borate buffer (pH 8.0). BSA–heparin–gold 5-nm colloid was centrifuged 30 min at 60,000 g and the pellet resuspended in a small volume of buffer. The procollagen and heparin–gold were mixed (25 μg of procollagen plus 0.25 ml of concentrated 5-nm heparin–gold), allowed to bind for 15 min at 25°C, and loaded into a Sepharose 4B column of 6.9-ml bed volume poured and equilibrated in 0.2 M (NH₄)₂CO₃ (pH 8.25). In some cases the reaction mixture was fixed with glutaraldehyde before loading on the column. The column was eluted at ∼5 ml/h and fractions of 0.2 ml were collected and analyzed by electrophoresis for protein, and by measuring A₅₂₀ to monitor heparin–gold content. Generally the first three fractions after the void volume were used for electron microscopy. Samples were prepared for electron microscopy as described by Margossian et al. (1991).

Electron Microscopy of Procollagen

Approximately 20-μg/ml solutions of complex were mixed with 0.25 vol of glycerol, sprayed on to freshly cleaved mica, dried in vacuo for 20 h, and rotary shadowed with platinum or tungsten by means of resistive or electron bombardment heating, respectively (Slayter, 1976, 1983). The estimated average metal-film mass thickness was 100 ng/cm² for tungsten and somewhat more for platinum. Samples were examined in the JEM 100CX electron microscope, using a top entry stage voltage with a 40-μm objective aperture, and at 100 kV acceleration voltage. Most micrographs were recorded at a magnification of 20,000× and enlarged photographically to 45,000× or 100,000×. High-resolution diffraction fields were also obtained, from the lightly shadowed specimen, by using matched annular condenser and objective apertures (Slayter, 1989, 1991).

Electron Microscopy of Fibrous Collagen

Reconstituted and native type I collagen fibrils were prepared as detailed above. Specimens of fibrous collagen were deposited on carbon-filmed grids, incubated at 4°C with heparin–gold for 30 min, washed with 0.1 M cacodylate buffer, pH 7.4, fixed 5 min with 0.25% glutaraldehyde in 0.1 M cacodylate buffer at pH 7.0, washed 3 × with water, and then stained with 1% uranyl acetate without thorough poststain washing (Slayter, 1983). Calibration was based on micrographs of grating- replica standards and of Indium-thione Olive crystals (Labaw, 1964).

Electron Microscopy of Frozen Sections

Blocks of rat tail tendon (0.5 mm) were fixed for 1 h in 0.1% glutaraldehyde in standard fixative (4% paraformaldehyde, 15% saturated in neutral picric acid). Tissue blocks were then immersed overnight at 4°C in PBS containing 2.3 M sucrose. These were maintained in a liquid nitrogen refrigerator until use, for a minimum of 24 h. Specimen blocks mounted on cutting stubs were removed and sections were obtained with thickness ∼60 nm. The methods used are essentially those of Tokoyasu and coworkers (Tokoyasu and Singer, 1976; Tokoyasu et al., 1983) as modified by Watkins.
et al. (1990). Thin sections were cut with a Reichert Ultracut Ultramicro-
tome fitted with an FC4D cryochamber (Vienna, Austria) at a temperature
range of -110 to -130°C. The sections were mounted on Formvar/carbon-
coated 200-mesh inch -1 grids and labeled as previously described (Geuze
et al., 1981). Sections were labeled by 30-min incubations with heparin-
gold at ~60 μg/ml, washed, counterstained with uranyl acetate, and em-
bedded in 1.25% methylcellulose as previously described (Watkins et
al., 1991). Sections were examined with a JEOL 100CX electron micro-
scope at 20,000× using a 40-μm objective aperture (JEOL U.S.A. Inc.,
Peabody, MA).

Results

Binding of Heparin to Different Forms
of Type I Collagen

The binding interaction of heparin with monomeric and
fibrillar collagen was studied using ACE. Type I collagen
was incorporated into ACE gels in the following forms:
procollagen type I (collagen monomers); reconstituted, acid-
solubilized type I collagen, and reconstituted, fibrillar type
I collagen. Electrophoresis of 125I-Tyr-heparin was con-
ducted through the collagen-containing lanes of ACE gels
and electrophoretograms are shown in Fig. 1 a. Retardation
coefficients (R) of the heparin migration fronts within each
ACE gel lane were measured, and are plotted versus protein
concentration (Fig. 1 b). From these curves the apparent av-
erage Kd’s of the binding between heparin and the collagen
preparations were derived. In two experiments, apparent
Kd’s (in nM) for heparin binding were: procollagen, 58 ±
2; acid soluble collagen, 79 ± 16; fibrillar collagen, 79 ±
14. These data indicate that the apparent affinities exhibited
by each of the collagen preparations for heparin are not
significantly different from each other. Furthermore, these
affinities are in the range of those exhibited by other heparin-
binding proteins for heparin (Lee and Lander, 1991; San An-
tonio et al., 1993).

Figure 1. (a) ACE analysis of the interactions between low Mr 125I-
Tyr-heparin and various forms of type I collagen. ACE gels were
constructed containing type I collagen in the procollagen, acid
solubilized, or reconstituted, fibrillar form. Collagens were present
in gel lanes at the concentration (nM) indicated beneath each lane,
with Mr’s taken to be 450 K for procollagen, and 300 K for the
other forms. Radiolabeled heparin was loaded into a sample
slot (located above each gel), and electrophoresis was conducted
towards the anode (located below each gel). Images of heparin
migration patterns within ACE gels were obtained using a phos-
phorimager (Molecular Dynamics). From these electrophoreto-
grams, the dissociation constant (Kd) can be estimated from the
protein concentration at which the heparin is half-shifted from be-
ing fully mobile at very low protein concentrations or between
protein-containing lanes, to being maximally retarded at high pro-
tein concentrations (Lee and Lander, 1991). (b) Calculation of
affinities of low Mr heparin for different forms of type I collagen.
Radiolabeled heparin was subjected to ACE analysis against the
various collagen preparations as illustrated in (a). Heparin retarda-
tion coefficients (R) within each collagen-containing lane were de-
termined (see Materials and Methods) and are plotted against pro-
tein concentration. Smooth curves represent nonlinear least-squares
fits to the equation $R = R_w/(1 + K_d/\text{protein}^2)$.
Visualization of Heparin-binding Sites on Type I Collagen Monomers and Fibrils

Mapping of Procollagen. To map the heparin-binding site on collagen monomers, electron microscopy was used to examine complexes between heparin-gold reagent and procollagen in solution. Rotary-shadowed preparations were made of the material obtained from the excluded volume of the gel filtration step used to separate unbound heparin-gold from that which is complexed with the procollagen (Fig. 2). We obtained micrographs of the complex material mixed with unbound procollagen. In the course of these experiments it was discovered that the fraction excluded at the column-void...
other hand, fraction 22 contained a much greater proportion termolecularly cross-linked collagen complexes associated glutaraldehyde, thus minimizing both intramolecular cross-
tations, buffered to neutrality. The fixation reaction was
hyde fixation, using a series of glutaraldehyde concentra-
tions. Procollagen molecules were measured for complexes ob-
constructed of overlapping collagen monomers. Thus, map-
ing experiments were undertaken with several forms of
fibrous collagen. Labeled fibrils were selected and scored for
the position of the heparin–gold particles in relation to the
67-nm repeat of native-type fibrils. Labeled fibrils were selected for
the position of heparin-gold attachment at ~300 nm from the center of the COOH-terminal head of procollagen. Preferred regions of heparin–gold
binding were found in all of the experiments (Fig. 4), and
the data are summarized in Table I. Most of the heparin–gold
is found at a site very near the tail end of the molecule. The peak at ~10–20 nm from the head, is somewhat suppressed at
the higher glutaraldehyde concentrations (Fig. 4, a and b),
but the other peak ~290 nm from the head (very near the
tail end), is always maintained (Fig. 4). Validity of these
results was supported by a control experiment in which bo-
vine serum albumin complexed with gold was used in mapping
experiments and failed to bind to procollagen (data not shown).

Mapping of Reconstituted Collagen Fibrils. Our finding
of one preferred heparin-binding site on procollagen mono-
mers led us to determine if heparin-binding sites are also
available in the type I collagen fibril, which is a structure
composed of overlapping collagen monomers. Thus, map-
ing experiments were undertaken with several forms of
fibrous collagen. Labeled fibrils were selected and scored for
the position of the heparin–gold particles in relation to the
67-nm repeat of native-type fibrils, measuring from the be-
"in the overlap zone (Fig. 4). The position of heparin-gold attachment at ~300 nm from the center of the COOH-terminal head of procollagen. Quantitative results are summarized in Table I.

To stabilize complexes, we subjected them to glutaralde-
hyde fixation, using a series of glutaraldehyde concentra-
tions, buffered to neutrality. The fixation reaction was
stopped by adding an excess of glycine to block unbound
glutaraldehyde, thus minimizing both intramolecular cross-
linking between procollagen molecules and background due
to possible nonspecific cross-linking between the hepa-
рин–gold and procollagen. Preferred regions of heparin–gold

Table I. Summary of Heparin–Gold–Procollagen
EM Mapping Data

| Experiment | GA Concentration | Histogram Peaks |
|------------|------------------|----------------|
| 1          | 0                | 195 maj 290 min |
| 2          | 0                | 20 maj        |
| 3          | .00002 Head/10 maj | 100 min      |
| 4          | .0001 Head/10 min | 280 maj      |
| 5          | .01              | -             |
| 6          | .01              | 290 maj       |

* Distance in nm from COOH terminus, or center of head, in procollagen, to center of heparin–gold marker.

GA, glutaraldehyde; maj, major peak; min, minor peak (collagen = 328 nm in same experiment).

EM Mapping Data

Figure 4. Histograms show mapping of the positions of heparin-gold on procollagen molecules in rotary-shadowed preparations of molecular complexes. Position of heparin-gold particles was measured from the center of the COOH-terminal globular head towards the NH₂ terminus of procollagen. Molecular complexes were fixed in: (a) 0.001%; (b) 0.0001%; (c) 0.00002% glutaraldehyde before column chromatography and rotary shadowing. In all cases there is a principle high frequency position of heparin-gold attachment at ~300 nm from the center of the COOH-terminal head of procollagen.
Figure 5. Composite correlating data obtained for the binding of heparin-gold to procollagen type I and to collagen type I fibrils. (a) Darkfield electron micrograph of tungsten rotary-shadowed preparation of procollagen showing globular COOH-terminal head on right, and position of putative heparin binding site (black arrow) as deduced by mapping data in Fig. 4. The location of the COOH-terminal end of tropocollagen and of the preferred heparin-binding site relative to the molecular overlap model of the collagen fibril is indicated by the dotted lines. (b) Diagram showing the arrangement of tropocollagen monomers within the collagen fibril, relative to the location of the overlap and gap zone fibril staining pattern. This molecular model was proposed by Chapman (1974). Tropocollagen molecules are shown as horizontal rods, and the polarity of all monomers in the fibril is indicated by N (NH$_2$ terminus) and C (COOH terminus) markings on one monomer. A diagram of the molecules within one 67-nm period (boxed area) is also shown expanded in Fig. 7. (c) Electron micrograph of glutaraldehyde-fixed, heparin-gold type I collagen fibril complexes, in fibrils visualized by uranyl acetate staining. Letters below micrograph show positions of positively stained fibril bands, following the accepted notation (Gross and Schmitt, 1948). Dotted lines between molecular model in b and the electron micrograph show corresponding overlap and gap zones. The location of heparin-gold particles relative to the molecular structure of the fibril as presented in Fig. 6 was measured within each 67-nm period, beginning at the center of the left border of the overlap zone (origin, arrow), and extending to the center of the right border of the gap zone. Heparin-gold particles appear as circular dark objects present mainly in the “a” bands region of the fibrils.

**Fibrils.** In another series of experiments the heparin-gold-binding site of native rat tail type I collagen was mapped before and after treatment of the collagen fibrils with chondroitinase. It was thought that chondroitinase treatment might uncover sites on collagen fibrils that are masked by endogenous proteoglycans such as decorin, thus making mapping of heparin-binding sites more definitive. In two experiments, the primary binding site was ~30 nm from the head. A second minor site at ~65 nm became somewhat more discrete after the chondroitinase treatment, but overall, chondroitinase treatment of fibrils had little effect on heparin-gold binding to fibrils. Table II shows histograms summarizing the data from these experiments. Since segment long spacing preparations of type I collagen are formed at low pH in acetic acid, and heparin-gold is unstable under these conditions, mapping experiments were found not possible to conduct on segment long spacing fibrils.

For the mapping experiments, the potential error in assigning positions on the basis of the heparin-gold label procedure is dependent upon several factors. First, the albumin-heparin-gold complex is ~10 nm in diameter. (The complex is visualized to be 10 nm in the metal-coated molecular preparations but only the 5-nm gold core is visualized in the fibrillar preparations.) Second, this marker could attach from the left, right, above, or below relative to the actual binding site. Third, some shrinkage upon dehydration (up to
10%) may be expected in procollagen. Finally, markers may detach slightly prior to or during fixation. However, the data show a number of rather narrow (10 nm, two histogram bars) histogram maxima peaks. It is thus concluded that all factors mentioned other than the marker size must be of lesser importance and suggest that the error is roughly the size of the marker.

**Discussion**

Here, we have explored the nature of heparin binding by type I collagen monomers and fibrils. In the first series of electron microscopy experiments, heparin binding to type I procollagen was studied, as procollagen remains in the monomeric state. Since procollagen retains its COOH-terminal globular head, the polarity of the molecules can be readily identified. Histograms of procollagen-heparin-gold mapping experiments include data from complexes that were unfixed as well as complexes fixed using three different glutaraldehyde concentrations. In all cases there is a principal high frequency position of heparin-gold attachment at ~30 nm from the beginning of the overlap zone toward the gap zone, in the “a” (see Figs. 5c and 7) bands region of the fibril.

![Figure 6. Histograms show mapping of the positions of heparin-gold on (a) reconstituted type I collagen fibrils; (b) native fibrils from homogenized rat tail tendons; (c) fibrils in frozen sections of native rat tail tendon. The location of heparin-gold particles relative to the molecular structure of the fibril (see Fig. 5) was measured within each 67-nm period, beginning at the center of the left border of the overlap zone, and extending to the center of the right border of the gap zone. In all cases there is a principal high frequency position of heparin-gold attachment at ~30 nm from the beginning of the overlap zone toward the gap zone, in the “a” (see Figs. 5c and 7) bands region of the fibril.](image)

**Table II. Summary of Heparin-Gold-Collagen Fibril EM Mapping Data**

| Experiment | Description                           | Histogram peak |
|------------|---------------------------------------|----------------|
| 7          | Reconstituted collagen                 | 30 nm          |
| 8          | Reconstituted collagen                 | 32 nm          |
| 9          | Reconstituted collagen                 | 30 nm          |
| 10         | Native collagen fibrils                | 30 nm          |
| 11         | Chondroitinase treated; native collagen fibrils | 30 nm          |
| 12         | Native fibrils                         | 27 nm          |
| 13         | Chondroitinase treated; native collagen fibrils | 27 nm          |
| 14         | Frozen section of nonfixed rat connective tissue | 27 nm          |

Fibrils was found near the interface between the overlap and gap region in the vicinity of the “a” bands as shown in Fig. 6. Experiments substantiated that this heparin-binding site occurs in the same position in reconstituted type I procollagen fibrils, as well as in those of native tissue, and in fibrils from homogenized rat tail tendons. The correlation between the results from the mapping of the heparin-binding site on procollagen with that of fibrils was facilitated using the overlap model presented by Chapman, 1974 (Fig. 5). In this model, the “a” bands contain elements of the NH2 terminus. Therefore, our finding that heparin attaches primarily to site(s) near the NH2 terminus of procollagen is compatible with the presence of a preferred binding site located near the division between the overlap and the gap zone in the fibrils, in the “a” bands region (see Fig. 5). All of the results with monomolecular complexes were obtained with type I procollagen obtained from chicken tendon fibroblasts. However, the experiments on fibrils used both native-type I collagen fibrils, and purified reconstituted native-type collagen fibrils from rat tail tendon. Given that the structural features of type I collagen are highly conserved, the comparison is most likely valid (Monson et al., 1982).

It is important to determine the level of structural organization of the collagen fibril required for high affinity heparin binding. Previous work has shown that monomeric tropocollagen binds heparin (Obrink, 1973), and that disrupting the triple helical conformation by melting of the collagen significantly reduces its heparin-binding affinity (Koda et al., 1985; Keller et al., 1986; San Antonio et al., 1992). In this report we have shown that type I procollagen and fibrillar type I collagen display similar binding affinities for low molecular weight heparin. Our finding that heparin-binding sites are available for the binding of heparin-gold not only in reconstituted collagen fibrils, but also in native collagen fibrils, which are complex structures composed of types I, III, and V collagen molecules, is also considered significant. In principle, one might expect fibrillar collagen to exhibit a higher apparent affinity than procollagen for heparin, since fibrils are potentially multivalent ligands. The fact that a higher apparent affinity was not seen suggests that any improvement in binding affinity due to multivalency is effectively offset in fibrils by a decrease in numbers of binding sites that are available. Such a decrease is expected, in fact, since a significant fraction of the collagen monomers in a...
fibril are thought to be buried beneath the surface, and presumably unavailable for binding.

Others have reported that when vertebrate collagenase was used to cleave type I collagen, the COOH-terminal fragment bound to a heparin affinity column more tightly than did the larger NH2-terminal fragment (Keller et al., 1986). This is contrary to our finding of a preferred binding site near the NH2-terminus, but there are ways in which this discrepancy may be explained. For example, it is possible that a cryptic heparin-binding site in the COOH-terminal region of type I collagen is made accessible for heparin binding only after collagenase treatment. Alternatively, differences in the characteristics (e.g., size, multivalency) of the heparin-gold agent used in the present study and the affinity matrix used by Keller et al. (1986), may give rise to differences in accessibility of heparin-binding sites to these two reagents.

It has been proposed that heparin-binding sequences in many proteins consist of clusters of three to five basic amino acids, interspersed with hydrophobic amino acids, and present within an α helical conformation (Cardin and Weintraub, 1989). When we searched the complete sequences of the αl and α2 chains of human type I procollagens (Kuivaniemi et al., 1988; Tromp et al., 1988) for such consensus sequences, none were found. Thus it appears that type I collagen uses novel types of binding sites to interact with heparin and HSPGs. As an approach to identifying the heparin-binding sites on type I collagen, we examined our data showing that the primary heparin-binding site on procollagen is located near the NH2-terminus of the molecule, and that within the fibril the preferred heparin-binding site falls in the "a" bands region (see Figs. 5 c and 7). We inspected the amino acid sequences of the αl and α2 chains of type I collagen in relation to the location of heparin binding we observed in the monomer and the fibril. In these regions of the collagen triple helices, we concentrated on locating amino acid sequences of net basic charge which might prove suitable as binding sites for the polyanion heparin. Interestingly, one highly basic sequence was found in the triple helix near the NH2-terminal end of procollagen, as well as within the "a" band region of the fibril, and is present in type I collagens of human, bovine, rat, and chick (Kuivaniemi et al., 1988; Hulmes et al., 1973; Tromp et al., 1988) (Fig. 7). This locus consists of nine basic amino acids, six of them contributed by the αl chains, and three by the α2 chain, as shown below:

αl: hlys-glyhis-arg-gly-phe
α2: hlys-glyile-arg-gly-his

In the αl chain of rat tropocollagen, this site includes amino acid residues 87-92 (Hulmes et al., 1973). Three of the basic amino acids within this sequence are hydroxylysine residues, which are involved in cross-links, and which sometimes function as dissacharide acceptor sites (Butler and Cunningham, 1965, 1966). We propose that this domain may function as a heparin-binding site in type I collagen. It should be stressed that there are other clusters of basic amino acids located in the collagen triple helices constituting the "a" band fibril region, but these all include fewer basic amino acids than the site discussed above, and often are present in close proximity to triple helical domains carrying strong negative charges (Fig. 7).

In the type I collagen monomer one other highly basic domain exists in the triple helix near the carboxy terminus (Fig. 7, arrow 7), and it also contains hydroxylysine residues involved in cross-links (Butler and Cunningham, 1966). The domain carries eight positive charges, and differs from the primary sequences of our proposed heparin-binding site, in both the composition and arrangement of some of its positively charged amino acids. It is possible that the weak heparin binding we observed near the COOH-terminal end of procollagen (Fig. 4, b and c) may be partially or wholly due to heparin binding to the COOH-terminal basic domain described above. In fact, this same site could be responsible for the heparin-binding affinity attributed to the COOH-terminal portion of type I collagen (Keller et al., 1986). The potential error in positioning the heparin-gold reagent relative to procollagen, however, makes it difficult at present to distinguish between binding of heparin-gold to the putative COOH-terminal heparin-binding sequence, as opposed to other basic domains near or on the globular head of procollagen.

It has been predicted that the proteoglycan-binding site should be in the "a" and "c2" band regions of the type I collagen fibril (see Figs. 5 c and 7), based on fibril staining patterns, as well as the concentration of unpaired positive charges of the combined rat and bovine sequences of the αl chains (Doyle et al., 1975). It is significant in the present study that we observed heparin binding mainly to the "a" bands of native type I collagen. Previous work showed, however, that in tissues the location of various types of proteoglycans (e.g., dermatan and keratan sulfate PGs) on type I collagen fibrils is not limited to the "a" bands region, but can also occur within the "c" and/or "e" bands regions (see Figs. 5 c and 7) of the 67-nm period (for review see Ruoslahti, 1988; Scott, 1988, 1991). Further work is necessary to determine if in vivo cell surface or extracellular matrix forms of HSPGs preferentially occupy the "a" bands region of type I collagen fibrils.

The relationships between the cell-attachment sites of type I collagen fibrils and heparin-binding sites are of great interest because heparin and syndecan-1 block cell adhesion to type I collagen in vitro (Koda et al., 1985; Piepkorn and Chapman, 1985). It is proposed that proteoglycans have this effect because they bind extracellular matrix molecules at sites that are adjacent to cell-attachment sites and thereby sterically hinder the binding of cell surface integrin receptors (Brennan et al., 1983).

The spatial relationship between the putative heparin-binding site and various potential cell-attachment sites on collagen fibrils are considered in Fig. 7. An average heparin molecule of 15,000 D is ~30-nm long. If collagen-binding sites on heparin consist of randomly distributed pentasaccharide sequences (as do antithrombin III-binding sites [Marcum and Rosenberg, 1989]), then a locus on heparin of ~2.5 nm could span the putative heparin-binding site on the collagen triple helix. In the collagen fibril, tropocollagen molecules are ~300-nm long, the intermolecular spacing between them is ~1.5 nm, and the length of one fibril repeat unit is ~67 nm (Brodsky and Eikenberry, 1982). Thus, heparin molecules would not be long enough to span the distance between two heparin-binding sites of adjacent 67-nm periods (see Figs. 5 and 7). Heparin molecules would, however, be long enough to bind to a heparin-binding site on one tropocollagen helix, and span the distance between this site, and certain cell-attachment domains on adjacent tropocollagen monomers (e.g., Fig. 7, sites marked by arrows 2 and

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protein matched nearly exactly with those of the rat, although it was the high degree of sequence homology between the collagens of acid (RGD) sequences. The model of Chapman (1974) for the overlap of the α1 chains of rat type I collagen within the fibril was used as a guide to align the α1 chains of the human molecule, based on the high degree of sequence homology between the collagens of these species. The distribution of charged residues of the human protein matched nearly exactly with those of the rat, although it was necessary to delete several nonhomologous residues from the human sequence to achieve alignment. The fibril portion shown is an expanded diagram of the molecules within the boxed area of Fig. 5 b, and includes one overlap and one gap zone, as indicated. Horizontal bars represent segments of five overlapping triple helical tropocollagen molecules, with the NH2 terminus (N) of one monomer located at the upper left of the diagram, and the COOH terminus (C) of another monomer at the lower right of the diagram; spacing between tropocollagen molecules is not to scale. Vertical bars represent total charge density at each residue unit along each triple helix, i.e., these values were determined by aligning the primary sequences of the human type I collagen α1 and α2 chains (Kuivaniemi et al., 1988; Tromp et al., 1988), and summing the charges of the three aligned residues across the chains of the triple helix. The location of the proposed heparin-binding domain is indicated (black arrow). Potential cell attachment sites and RGD sequences include DGEA (arrow 1) (Staatz et al., 1990, 1991) of the α1 chains of human, chicken, bovine, and rat; RGDA of the α1 chains of the human (arrow 2); RGDT of the α2 chain of the chicken but not human protein (arrow 3) (Dedhar et al., 1987; Pignatelli and Bodmer, 1988); RGDG of the α2 chain of the human (arrow 4); RGDM of the α2 chain of the human (arrow 5); and RGDK of the α1 and α2 chains of the human (arrow 6) (see Hulmes et al., 1973; Bernard et al., 1983; Kuivaniemi et al., 1988; Tromp et al., 1988 for all sequences). Basic domain near COOH terminus showing sequence homology to proposed heparin-binding site (see Discussion) is indicated (arrow 7). Letters and bars at bottom of figure 3), thereby potentially influencing cell–collagen interactions. The precise in vivo role of HSPGs as regulators of integrin receptor–collagen interactions will remain uncertain, however, until the locations of available integrin-binding sites on native type I collagen fibrils have been mapped.

It is of interest to speculate on the in vivo role of the heparin-binding site of type I collagen. Possibly, this site functions to join extracellular matrix PGs and collagen fibrils, to give structural support to certain connective tissues, or to control aspects of collagen fibrillogenesis (e.g., see Ruoslahti and Engvall, 1980; Vogel et al., 1984; Ruoslahti and Pierschbacher, 1987; Scott and Parry, 1992). Alternatively, the heparin-binding site of type I collagen may function in cell adhesion, as a primary matrix receptor for cell surface HSPGs (Koda et al., 1985; Sanderson et al., 1992). For example, via this binding site, cell surface HSPGs may act as coreceptors, by bringing extracellular ligands (e.g., collagen) together with high affinity cell surface receptors for these ligands (e.g., integrins) (see Bernfield et al., 1992 for review). Thus, given the abundance of HSPGs and type I collagen fibrils in most tissues, it is likely that the heparin-binding site on type I collagen participates in key physiological processes such as cell adhesion, migration, or cell-directed assembly or restructuring of the extracellular matrix.

We gratefully acknowledge the contributions of Dr. Bjorn R. Olsen of Harvard Medical School, for his expert advice on collagen isolation and biochemistry; Dr. Gerald Tromp of Thomas Jefferson University, for supplying sequence data on human type I collagen; Dr. Yuhui Xu, for providing micrographs of frozen sections of connective tissue; Ms. Elizabeth Gunning for expert assistance in data analysis and manuscript preparation; and Dr. Michael E. Oettlinger and Dr. Marla J. Steinbeck of Harvard Medical School for critical review of the manuscript.

This work was supported by National Institutes of Health grant HL 33041 to H. Slayter, HL 17747 to M. J. Karnovsky, NS 26862 to A. D. Lander, and an Epply Foundation grant to J. D. San Antonio.

Received for publication 11 October 1993 and in revised form 10 February 1994.

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