Interaction Between Amino Propeptides of Type XI Procollagen 1 Chains

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Received for publication, September 16, 2003, and in revised form, December 18, 2003
Published, JBC Papers in Press, December 29, 2003, DOI 10.1074/jbc.M310291200

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Type XI collagen is a quantitatively minor yet essential constituent of the cartilage extracellular matrix. The amino propeptide of the α1 chain remains attached to the rest of the molecule for a longer period of time after synthesis than the other amino propeptides of type XI collagen and has been localized to the surface of thin collagen fibrils. Yeast two-hybrid system was used to demonstrate that a homodimer of α1(XI) amino propeptide (α1(XI)Npp) could form in vivo. Interaction was also confirmed using multi-angle laser light scattering, detecting an absolute weight average molar mass ranging from the size of a nonamer to the size of dimer (25,000–50,000 g/mol), respectively. Binding was shown to be saturable by ELISA. An interaction between recombinant α1(XI)Npp and the endogenous α1(XI)Npp was observed, and specificity for α1(XI)Npp but not α2(XI)Npp was demonstrated by co-precipitation. The interaction between the recombinant form of α1(XI)Npp and the endogenous α1(XI)Npp resulted in a stable association during the regeneration of cartilage extracellular matrix by fetal bovine chondrocytes maintained in pellet culture, generating a protein that migrated with an apparent molecular mass of 50–60 kDa on an SDS-polyacrylamide gel.

Type XI collagen is a constituent of the extracellular matrix of cartilage. It belongs to the family of fibrillar collagens, which includes types I, II, III, V, and XI (1). As with all of the fibrillar collagens, type XI contains a 300-nm uninterrupted triple helical domain (2). Type XI collagen forms the heterotypic collagen fibrils characteristic of cartilage along with types II and the nonfibrillar type IX collagen (3). Although minor in quantity, type XI collagen is thought to play a critical role in collagen fibril assembly and extracellular matrix organization in early development as demonstrated by the chondrodystrophic mouse (cketocho) (4). Fibers of developing cartilage are of uniform 20 nm in diameter and oriented more randomly than fibrils of adult articular cartilage.

Collagen fibril assembly in embryonic cartilage is known to rely upon the ratio of the collagenous constituents, types II and XI (5). Initially, type XI collagen may function to nucleate the formation of new collagen fibrils (6). The role of type XI collagen in limitation of lateral growth of collagen fibrils is well established, although the molecular mechanism is not fully understood. Type XI collagen may also play a role in the maturation of thin fibrils into the larger diameter fibrils of the territorial zone of cartilage matrix, either by regulation of the accretion of more collagen molecules onto the surface of thin fibrils or by mediating the fusion of thin fibrils.

Type XI collagen is initially synthesized as a procollagen, which is subsequently posttranslationally processed at both the amino and carboxyl termini (7). The amino-terminal processing site was identified 7 amino acids from the variable region (8, 9). Recently, type XI α1 pro-collagen was identified as a substrate for the enzyme bone morphogenetic protein 1 (BMP-1) (10). The carboxyl propeptide of the α1 chain of type XI collagen is homologous to the other fibrillar collagens, whereas the amino propeptide of type XI α1 pro-collagen is markedly different. In addition to the characteristic minor triple helix, the α1 chain of XI contains a large non-triple helical domain at the amino terminus.

The α1 chain of type XI collagen exists as a set of isomers that arise by alternative splicing of the primary RNA transcript (11). The portion encoding the variable region gives rise to biochemically unique domains. However, common to all isomers is the amino propeptide, the distal globular 233 amino acids, which is ultimately posttranslationally removed during the process of extracellular matrix assembly and fibril formation. Proteolytic removal of the amino propeptide may rely on the action of BMP-1, as it does in the case of α1(V) collagen chain (12).

The amino propeptide domain is structurally related to other amino propeptides including that of the α2(XI) and α1(V) collagen chains. The position of four cysteine residues are conserved, and sequence analysis predicts a conserved secondary structural motif rich in β-strand (13). This prediction has been supported by analysis of circular dichroism spectra, demonstrating a relatively high percentage β-strand and a markedly low percentage α helix (14). The dimensions of the globular amino propeptide domain of α1(XI) collagen chain has been estimated by rotary shadowing to be at least 8 nm in diameter. This domain has been localized to the surface of collagen fibrils by immunoelectron microscopy (15).

Although the amino propeptides are ultimately removed from the collagen triple helical molecule by proteolytic processing, the rate and extent of removal vary. Although the processing of α1(XI) collagen chain is relatively slow, thus achieving 50% completion by 18 h after synthesis in pulse-chase experi-

* This work was supported by the Arthritis Foundation, the Gerlinger Foundation, National Institutes of Health Grants R01AR47985, K02AR48672, and P20RR18454-Biomedical Research Infrastructure Network for Idaho (BRIN). The costs of publication of this article were defrayed in part by the payment of page charges. This article must therefore be hereby marked ‘advertisement’ in accordance with 18 U.S.C. Section 1734 solely to indicate this fact.

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EXPERIMENTAL PROCEDURES

Synthesis and Purification of Recombinant Rat a1(XI)Npp Domain—A cDNA fragment encoding the amino propeptide of a1 chain of type XI collagen was inserted into expression vector pET11a (Stratagene). BL21 DE3 transformants were screened for expression of recombinant protein on SDS-polyacrylamide gels and subsequent transfer to polyvinylidene difluoride membrane for detection of recombinant proteins by nickel affinity blot using nickel-NTA conjugated to alkaline phosphatase (Qiagen). Cells from transformant cultures were collected by centrifugation, and proteins were solubilized using bacterial protein extraction reagent (Pierce) with 1 unit of DNase I per 10 ml of cell suspension. This extraction solubilized the majority of cellular proteins, leaving recombinant a1(XI) amino propeptide (a1(XI)Npp) insoluble in the inclusion bodies. Protein from inclusion bodies was solubilized and unfolded in 20 mM Tris-HCl, pH 7.5, with freshly prepared 8M urea, and 14 mM mercaptoethanol at a protein concentration of ~1 mg/ml. Refolding was initiated by diluting solubilized unfolded protein drop-wise and with stirring into 100 volumes of 20 mM Tris-HCl, pH 7.5, containing 100 mM KCl, 20% glycerol, and 2 mM urea at 4 °C. Incubation at 4 °C was continued for 16 h. Refolded protein was concentrated by ultrafiltration using a Centriprep filtration unit (Amicon). Insoluble material, if present, was removed by centrifugation. Refolded protein was further purified by chelation chromatography on a HiTrap nickel column (Amersham Biosciences) (Fig. 1). Disulfide-bonded cysteine pairs were verified by tryptic digest followed by protein sequencing of those bands present in the absence of 10 mM DTT but absent when the sample was treated with reducing agent as described previously (17). Refolded protein was characterized by circular dichroism and compared with native protein isolated from tissue (17). Npp without a His6 tag was generated by removing the carboxyl portion of a recombinant a1(XI) amino-terminal domain containing the variable region (a1(XI)NTD), which contains a functional BMP-1 proteolytic processing site (10). Recombinant protein was incubated in the presence of purified recombinant human BMP-1 (FibroGen, Inc.) as described previously (10).

Yeast Two-hybrid System—A protein-protein interaction was tested in vivo using Matchmaker GAL4 (Clontech, Inc.) yeast two-hybrid system. cDNA encoding a1(XI)Npp was obtained by reverse transcription-PCR using the following primers: 5′ - cca tgg tgt aca tgc aag ctt tag at-3′ and 5′ - aag ccc ttt gat ctc atc atg cgt tag at-3′. This fragment was used to make two different gene fusions: one with the DNA binding domain of GAL4 and the other with the transcriptional activation domain of GAL4 in the plasmids pGBK7 and pGAD7, respectively.

Yeast host strain AH109 was transformed using the lithium acetate method (17). Selection of positive transformants was carried out by plating on SD minimal medium agar plates lacking the nutrient leucine and tryptophan to select for the plasmids derived from pGBK7 and pGAD7, respectively. Plates lacking both nutrients were used to select for double transformants.

To select for colonies in which the two-hybrid proteins interact, double transformants were plated on SD minimal medium lacking tryptophan, leucine, and histidine. Protein-protein interaction was verified by determining the expression of β-galactosidase activity by colony-lift filter assay and by determining the level of expression in liquid culture using chloroform red-β-galactosidase as substrate (Clontech).

Weight Average Molecular Weight Determination by Light Scattering—Recombinant a1(XI)Npp was applied to a TSK 3000SW size exclusion column at 10 °C in the following buffer: 30 mM Na2HPO4, 45 mM KH2PO4, pH 6.6, containing 100 mM KCl. The eluting peak was analyzed using a refractive index detector, a UV monitor, and by multi-angle laser light scatter (DAWN, Wyatt Technology, Santa Barbara, CA) to determine the absolute molecular weight of a1(XI)Npp in solution. A vertically polarized laser beam was passed through the flow cell containing the sample. The scattered light was simultaneously detected by 18 separate detectors at different angles. The intensity of the scattered light was digitized and transmitted to a computer for analysis by software provided by the manufacturer. The weight-average molar mass was determined from the Rayleigh ratio of scattered light, molecular concentration in grams/milliliters, and the specific refractive index increment of the dissolved molecules (dn/dc). The concentration was determined from the refractive index at each slice of the chromatogram and also from the absorbance at 280 nm and the 0.1% extinction coefficient, calculated for recombinant rat a1(XI)Npp to be 0.548 assuming all cysteine residues exist as half-cystines. The dn/dc value used was 0.185. Recombinant a1(XI)Npp was analyzed under three salt conditions: 75, 175, and 500 mM with respect to additive concentration of KCl and buffer components. Mol- lars was determined for individual slices of data across the eluting size exclusion chromatographic peak and plotted as a function of elution volume. Molecular weight at the leading and trailing edges of the peak was compared to determine whether recombinant a1(XI)Npp multimerized under the buffer conditions of the column.

Antibodies—A peptide coding for the human a1(XI)Npp was used to make the a1(XI)Npp antibody that showed specificity for the endogenous a1(XI)Npp but not the recombinant a1(XI)Npp. Preparation and characterization of this antibody was reported previously (18). The ability of the a1(XI)Npp antibody to discriminate between endogenous and recombinant a1(XI)Npp was demonstrated by immunoblot and nickel affinity blots (Fig. 6). Samples of increasing amounts of recombinant a1(XI)Npp were loaded onto an SDS gel and subsequently transferred to polyvinylidene difluoride membrane. Nickel-NTA agarose was used to determine whether recombinant a1(XI)Npp from a sample that was three orders of magnitude more dilute than the concentration needed for detection by the polyclonal antibody to a1(XI)Npp. Generation of the a2(XI)Npp antibody was carried out using the sequence RERPQRQPSHRTQ from bovine sequence (18) as the immunizing peptide. Antibody to the variable region of the recombinant a1(XI) amino-terminal domain was described previously (19).

ELISA Assay—Recombinant a1(XI)Npp was adsorbed to a microtiter plate in 96-well format at room temperature. To block unoccupied sites on plastic surface, wells were incubated with buffered skim milk powder solution for 1 h at room temperature. Recombinant a1(XI)NTD was adsorbed to a microtiter plate in 96-well format at room temperature. Antibody to the unique variable region was used to detect interaction. Absorbance at 405 nm was plotted as a function of the concentration of bound a1(XI)NTD with variable region detected. This antibody did not recognize a1(XI)Npp. Nonspecific binding of the primary antibody was tested in the absence of antigen. Nonspecific binding of the secondary antibody was tested in the absence of primary antibody. Bovine serum albumin was bound to the 96-well plate as a control to detect nonspecific binding of a1(XI)NTD. Data was subjected to Scatchard analysis to determine KE and Bmax.

Co-precipitation of Recombinant and Endogenous a1(XI)Npp Domains—Recombinant a1(XI)Npp, synthesized with six histidine residues at the carboxy terminus of the cDNA encoding a1(XI)Npp, was reacted with nickel-NTA agarose (Qiagen) to investigate the specificity of the interaction between a1(XI)Npp domains. Protein was extracted from bovine chondrocyte pellet cultures using 50 mM Tris-HCl, pH 7.5, 1 M NaCl, and 5 mM EDTA. No denaturants such as guanidine hydrochloride or urea were present during the extraction. Salt concentration was re-adjusted to 150 mM by dialysis prior to incubation with a1(XI)Npp-bound nickel-agarose. Material that bound to the recombinant a1(XI)Npp was separated from that which did not bind by centrifugation. Samples were analyzed by SDS-polyacrylamide gel electrophoresis followed by immuno-blot using an antibody that recognized a1(XI)Npp and one that recognized a very similar domain, a2(XI)Npp.

Pellet Culture of Bovine Chondrocytes—Bovine chondrocytes were maintained in pellet culture (20). Where indicated in the text, β-aminopropionitrile fumarate was included in pellet culture medium at a final concentration of 64 μg/ml, putrescine was included at a final concentration of 2 mM, AEBFS (4-(2-aminoethyl)-benzene sulfonyl-
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FIG. 1. Expression of recombinant rat α1(XI)Npp. A, schematic representation of the amino-terminal domain of a type XI collagen molecule. The Npp of the α1(XI) chain is shown as a terminal globular domain attached to the minor helix (mh) via an extended variable region (vr). The α2(XI)Npp is absent from this schematic to represent the rapid proteolytic processing of this chain. Between the minor helix and the major triple helix (TH) lies the amino telopeptide (tp). Black bar represents location of antibody recognition site. B, extracts of E. coli cells grown in the presence of IPTG were resolved by SDS-PAGE on a 12% polyacrylamide gel stained with Coomassie Blue. Lane 1, molecular weight markers; lane 2, mid-log culture grown in the absence of IPTG; lane 3, mid-log culture grown in the presence of IPTG; lane 4, insoluble protein after extraction of E. coli cell pellet; lane 5, soluble protein after protein extraction; lane 6, flow-through from nickel-affinity column; lane 7, column wash; lanes 8 and 9, elution of purified, refolded recombinant protein; lane 10, recombinant protein markers.

RESULTS

Recombinant Rat α1(XI)Npp Domain—A 682-bp fragment encoding α1(XI)Npp was obtained by reverse transcription-PCR and inserted into the pET11a expression vector. Recombinant α1(XI)Npp was expressed as a fusion protein with gene-10 gene product of bacteriophage T7, resulting in the addition of amino acids MAS at the amino terminus of the protein. Protein was expressed by BL21(DE3) Escherichia coli cells upon isopropyl-1-thio-β-d-galactopyranoside induction. The DNA sequence of the insert was determined by automated fluorescence DNA sequencing. The terminal methionine was removed in the bacterial expression system as verified by LCQ™MS/MS. The sequence of interest proceeded from the first proline after the signal peptide processing site of α1(XI) and continued to the glutamine one amino acid before the amino propeptide processing site. Codons for six histidine residues were included in the downstream PCR primer immediately preceding a stop codon. The penultimate amino acid, alanine, was changed conservatively to a valine to allow the antibody to discriminate between endogenous bovine α1(XI)Npp and recombinant rat α1(XI)Npp. In addition, this change modified the sequence at the proteolytic processing site of α1(XI) collagen, reducing the chance that the histidine tag would be removed from the recombinant protein. Recombinant α1(XI)Npp with a His<sub>6</sub> tag had a molecular weight of 24,826 g/mol and a theoretical pI of 6.91. Recombinant α1(XI)Npp was recovered from inclusion bodies. After unfolding and refolding, recombinant α1(XI)Npp was purified by nickel-chelation chromatography (Fig. 1). Purified bacterial recombinant α1(XI)Npp co-migrated with human embryonic kidney 293 recombinant α1(XI)Npp by SDS-polyacrylamide gel electrophoresis (data not shown). Unfolded and refolded α1(XI)Npp was previously characterized by spectropolarimetry, electron microscopy, and LCQ-MS/MS (17).

Detection of Interaction Using Yeast Two-hybrid System—Expression vectors encoding fusion proteins between the DNA binding domain of GAL4 and α1(XI)Npp and between the transcriptional activation domain of GAL4 and α1(XI)Npp were constructed. Cells cotransfected with both constructs were able to grow in the absence of histidine (Fig. 2), indicating an interaction between the α1(XI)Npp domain portion of each of the fusion proteins. Interaction was also demonstrated by detection of β-galactosidase activity (Fig. 2C). The level of β-galactosidase activity in double transformants was ~10-fold higher than background.

Size Exclusion Chromatography and Multi-angle Laser Light Scatter—At physiological ionic strength (175 mM), the weight average molar mass varied from the molecular mass of a monomer (25,000 g/mol) at the trailing edge of the size exclusion chromatography peak to 40,000 g/mol, an average molar mass lying between that of a dimer (50,000 g/mol) and a monomer at the leading edge of the peak (Fig. 3). Interaction was favored by low ionic strength (75 mM) and disrupted by high ionic strength (500 mM). Maximum molar weight observed corresponded to a dimer with no indication of higher multimerization. Multimerization was inferred by an increase in molar mass of purified recombinant α1(XI)Npp.

ELISA—Binding was shown to be saturable by ELISA (Fig. 4) in which recombinant α1(XI)Npp was bound to the plate and a tagged recombinant isoform of α1(XI) amino-terminal domain was allowed to interact with the α1(XI)Npp bound to the plate. Antibody to the unique region of the alternate isoform was used to detect interaction. This antibody did not recognize α1(XI)Npp. The dissociation constant was determined to be on the order of 0.13 μM.

Specificity of α1(XI)Npp Interaction: Co-precipitation of Endogenous and Recombinant α1(XI)Npp—Bovine chondrocytes maintained in pellet culture were used as a source of cartilage extracellular matrix proteins. Extracts were prepared from 10-day pellets using 1M NaCl in Tris buffer in the absence of denaturants. These extracts were incubated with recombinant α1(XI)Npp bound to agarose beads. Samples were analyzed by SDS-polyacrylamide gel electrophoresis and subsequent immunoblot (Fig. 5). As a control, endogenous α2(XI)Npp was analyzed for its ability to bind to recombinant α1(XI)Npp under the same conditions. Using an α2(XI)-specific antibody, α2(XI)Npp was found primarily in the free fraction, whereas bovine α1(XI)Npp bound to the recombinant α1(XI)Npp-agarose beads (Fig. 5).

Specificity of Antibody for the Endogenous α1(XI)Npp—The ability of the α1(XI)Npp antibody to discriminate between endogenous and recombinant α1(XI)Npp was demonstrated by immunoblot and nickel affinity blot (Fig. 6). Samples of increasing amounts of recombinant α1(XI)Npp were loaded onto an SDS gel and subsequently transferred to polyvinylidene difluoride membrane. Nickel-NTA-alkaline phosphatase conjugate was used to detect recombinant protein from a sample that was three orders of magnitude more dilute than the concentration needed for detection by the polyclonal antibody to α1(XI)Npp. Location of the epitope site within the amino propeptide and differences in amino acid sequence among the peptide antigen, endogenous bovine sequence, and the sequence within the recombinant protein are demonstrated in Fig. 6C.

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Interaction in Chondrocyte Pellet Culture System Resulted in a Stable Association—Recombinant α1(XI)Npp was converted from a protein that migrated as an ~30-kDa protein to one that had an apparent molecular mass of ~60 kDa under denaturing and reducing conditions when incubated for 24 h with bovine chondrocytes maintained in pellet culture. Both the 30- and 60-kDa bands were detectable via the His6 tag by Nickel affinity blot (Fig. 7). Using the α1(XI)Npp antibody that can discriminate between recombinant and endogenous bovine α1(XI)Npp, the 60-kDa band was also recognized by this antibody, suggesting the formation of a dimer of α1(XI)Npp monomers, one contributed by the endogenous bovine α1(XI) and the other by the added recombinant α1(XI)Npp. In addition, a band with an apparent molecular mass of 35 kDa was identified as immunologically related to endogenous α1(XI)Npp. The nature of this band is currently unknown. Stability of the 60-kDa band...
Fig. 4. **ELISA assay.** Binding was shown to be saturable by ELISA assay in which recombinant α1(XI)Npp was bound to the plate and tagged recombinant α1(XI) amino-terminal domain was allowed to interact with the bound α1(XI)Npp. Antibody to the unique tag region was used to detect interaction. Absorbance at 405 nm was plotted as a function of tagged α1(XI) concentration (■). This antibody did not recognize α1(XI)Npp. Inset depicts the results of Scatchard analysis of binding. Bovine serum albumin was used as a control for nonspecific interaction (✖). Error bars indicate ± S.E.

Fig. 5. Specificity of α1(XI)Npp interaction by co-precipitation of endogenous α1(XI)Npp with recombinant α1(XI)Npp-nickel-agarose. Proteins were extracted from chondrocyte pellet cultures in 1 M NaCl-containing buffer. Buffer conditions were changed by dialysis to reduce the salt concentration to 150 mM NaCl, and extract was incubated with α1(XI)Npp-bound nickel-agarose beads. Proteins were resolved by SDS-PAGE 12% polyacrylamide gel and analyzed by immunoblot using an antibody to α1(XI)Npp and an antibody to α2(XI)Npp. Lanes 1 and 3, material not bound to α1(XI)Npp-bound nickel-agarose beads. Lanes 2 and 4, material bound to α1(XI)Npp-bound nickel-agarose beads. Lanes 1 and 2, immunoblot with α2(XI)Npp antibody. Lanes 3 and 4, immunoblot with α1(XI)Npp antibody. Lane 5, molecular weight markers. Arrows on left-hand side indicate position of the α1(XI)Npp and α2(XI)Npp.

was investigated by subjecting samples to prolonged incubation in the presence of 250 mM imidazole, EDTA, or β-mercaptoethanol at 90°C with no change observed in migration on SDS gels. The generation of the 60-kDa band during 24 h in bovine pellet culture was not inhibited by inclusion of β-aminopropionitrile fumarate to inhibit lysine-derived covalent cross-linking or by inclusion of putrescine to inhibit transglutaminase in the culture (data not shown). To rule out a possible interaction mediated by the His₉ tag present on the recombinant protein, recombinant α1(XI)Npp was generated without a His₉ tag, by removing the carboxyl domain from the recombinant α1(XI) amino-terminal domain isofom by enzymatic proteolysis with BMP-1 (Fig. 7, lanes 6 and 7). The resulting recombinant α1(XI)Npp was included in chondrocyte cell culture with similar results (Fig. 7).

**DISCUSSION**

The α1 chain of type XI collagen exists as a set of isoforms that arise by alternative splicing of the primary RNA transcript. The portion encoding the variable region gives rise to biochemically unique domains. However, common to all isoforms is the amino propeptide, the distal globular 233 amino...
for any of the other fibrillar collagens and no such interactions have been described between recombinant domains. In the case of types IV and X, the interactions were hydrophobic in nature rather than due to covalent bond formation. In the case of the type XI a1 Npp dimer, the stable interaction takes place only in the presence of living cells because incubation of purified Npp does not result in the appearance of higher molecular weight domain on subsequent SDS gels, neither does the incubation of mixtures of recombinant isoforms of a1(XI) amino-terminal domain that contains any of the possible variable domains adjacent to the Npp domain (data not shown). This interaction may because of hydrophobic interaction or may be the result of covalent bond formation between the two Npp subunits. Data from the light scattering experiments do not support a hydrophobic interaction, because a decrease in average molecular weight was observed with increasing ionic strength of the buffer. If the stable interaction is the result of the formation of a covalent bond, it is unknown what enzyme is responsible at this time.

Type XI collagen serves primarily a regulatory function in collagen fibrillogenesis. The interaction between a1(XI)Npp domains demonstrated in this study may be coupled to enzymatic events including covalent cross-linking of amino propeptide domains and proteolytic cleavage to remove these domains from collagen fibrils. Such an interaction could contribute to our understanding of 1) initial nucleation of collagen fibrils, 2) lateral fusion of preexisting thin fibrils, and 3) increase in diameter because of accretion of individual collagen molecules to the surface of a growing collagen fibril.

The role of type XI collagen in limitation of lateral growth of collagen fibrils is well established, although the molecular mechanism is not fully understood. The a1(XI)Npp has been localized to the surface of thin collagen fibrils, it is relatively long-lived at the surface, and it is clear from the molecular dimensions that the a1(XI)Npp, the variable region, and the minor helix would not be accommodated within the gap region and would therefore sterically hinder further accretion of type II collagen molecules to the surface of a growing fibril. An interaction between Npp domains that favor proteolytic removal of globular domains from the surface may facilitate lateral growth of collagen fibrils.

Acknowledgments—We thank Dr. Lin Randall and Traci Topping for assistance with the size exclusion chromatography/multi-angle light scattering experiments.

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